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(21) Application number: 92100816.5

(22) Date of filing: 18.01.1992

(12)

(54) Substituted 1,2,3,4-tetrahydrocyclopent[b]indoles,

1,2,3,3a,4,8a-hexahydrocyclopent[b]indoles and related compounds, intermediates and a process for the preparation thereof and their use as medicaments

Substituierte 1,2,3,4-Tetrahydrocyclopent[b]indole, 1,2,3,3a,4,8a-Hexahydrocyclopent[b]indole und verwandte Verbindungen. Zwischenprodukte und ein Verfahren zur Herstellung derselben und ihre Verwendung als Medikamente

1,2,3,4-Tétrahydrocyclopent[b]indoles, 1,2,3,3a,4,8a-hexahydrocyclopent[b]indoles substitués et composés apparentés, intermédiaires et un procédé pour leur préparation et leur utilisation comme médicaments

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Description

The present invention relates to compounds of the formula,

$$R_4$$

$$X$$

$$R_3$$

$$R_2$$

$$R_2$$

$$R_2$$

where

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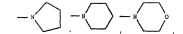
n is 2, 3, 4 or 5;

X is hydrogen, C_1 - C_2 -eilkilly, C_2 - C_2 -eilkovy, hydroxy, halogen, trifluoromethyl or nitro, R_1 is hydrogen, C_1 - C_2 -alkyl, C_2 - C_2 -cycloalkilly, C_2 - C_2 -alkyl, C_2 - C_2 -alkyl, C_2 - C_2 -cycloalkyl, wherein the phenyl group is substituted with 0, 1 or 2 substituents, each of which being independently, C_2 - C_2 -alkov, C_2 - C_2 -alkov, hologen, trifluoromethyl, hydroxy or nitro.

the group "Alk" signifying a divalent C_1 - C_6 -alkylene group, and Y signifying hydrogen, C_1 - C_6 -alkyl, phenyl or phenyl- C_2 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above; R_2 is hydrogen, C_1 - C_6 -alkyl, formyl, C_1 - C_6 -alkylcarbonyl, benzyloxycarbonyl or C_1 - C_6 -alkylaminocarbonyl; or alternatively, the group



as a whole is



$$NH - N - C_1 \cdot C_6$$
-alkyl

or

$$N-C_1-C_6$$
 -alkylphenyl

35 wherein the phenyl group may be substituted as indicated above,

 R_0 is hydrogen, C_1 - C_6 -alkyl, phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above; C_1 - C_6 -alkylicarbonyl or C_1 - C_6 -alkoxycarbonyl; R_4 is hydrogen. OH.

or

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wherein

 R_g is C_1 - C_6 -alkyl, C_2 - C_6 -alkynyl, C_3 - C_7 -cycloalkyl, C_3 - C_7 -cycloalkyl- C_1 - C_6 -alkyl, phenyl, phenyl- C_1 - C_6 -alkyl or phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above, and R_g is hydrogen, C_1 - C_6 -alkyl, phenyl or phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above, or alternatively the group

$$-N$$
 R_5

as a whole is

$$-N$$
 NH $-N$ $N-C_1-C_6$ -alkyl $-N$ N -phenyl

40 or

wherein the phenyl group may be substituted as indicated above,

and

R₇ is C₁-C_e-alkyl, phenyl or phenyl-C₁-C_e-alkyl, wherein the phenyl group may be substituted as indicated above:

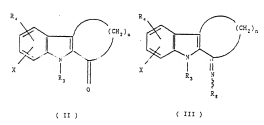
with the proviso that R₄ is not hydrogen or hydroxy, when n is 4 or 5; or a pharmaceutically acceptable acid addition salt thereof.

which compounds are useful for alleviating various memory dysfunctions characterized by a cholinergic deficit such as Alzheimer's disease. Compounds I of this invention also inhibit monoamine oxidase and/or act at central c₂-adrenergic receptors, and hence are useful as antidepressants.

Also included within the scope of this invention are compounds of Formula II, where R₃, R₄, X and n are as previously defined, which are useful as direct precursors to the target compounds of this invention.

Also included within the scope of this invention are compounds of Formula III, where R₆ is hydroxy, amino-C₁-C₆alkoxy, C₁-C₆-alkyl, C₂-C₇-cycloalkyl, C₂-C₇-cycloalkyl, phenyl-C₁-C₆-alkyl or phenyl-C₃-C₇-cycloalkyl, wherein the phenyl orous may be substituted as indicated above:

C₁-C₆-alkylcarbonyloxy or C₁-C₆-aminocarbonyloxy, which are useful for alleviating various memory dysfunctions characterized by a cholinergic delicit such as Alzheimer's disease. Compounds III of this invention also inhibit monoamine oxidase and/or act as presynaptic «-aefonoratic receptor antaconists, and hence are useful as anticlopressants.



Unless otherwise stated or indicated, the following definitions shall apply throughout the specification and the appended claims.

The term loweralkyl shall mean a straight or branched alkyl group having from 1 to 6 carbon atoms. Examples of said loweralkyl include methyl, eithyl, n-propyl, iso-propyl, n-butyl, iso-butyl, sec-butyl, t-butyl and straight- and branched-chain pentyl and haxyl.

The term halogen shall mean fluorine, chlorine, bromine or jodine.

Throughout the specification and the appended claims, a given chemical formula or name shall encompass all stereo and tautomeric isomers where such isomers exist.

The compounds of this invention are prepared by utilizing one or more of the synthetic steps described below.

Throughout the description of the synthetic steps, the notations n, X, Y and R_1 through R_0 shall have the respective meanings given above unless otherwise stated or indicated.

STEP A:

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A compound of Formula IV, where $\rm R_{\rm g}$ is hydrogen or -OCH $_{\rm g}$, is allowed to cyclize to afford a compound of Formula V. This reaction is typically conducted in aqueous sulfuric acid at a temperature of 25 to 150°C.

(IV)

(V)

STEP B:

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Compound V is allowed to react with a sulfate compound of the formula, $(R_{10}O)_2SO_2$, where R_{10} is $C_1 \cdot C_6$ -alkyl or phenyl- $C_1 \cdot C_6$ -alkyl, wherein the phenyl group may be substituted as indicated above, in a routine manner known to the at to afford a compound of Formula VI.

Alternatively, compound V is allowed to react with a halide compound of the formula R₁₀ - Hal, where R₁₀ is as defined above, in a routine manner known to the art, to afford a compound of Formula VI.

$$(V) + (R_{10}O)_2SO_2$$
 X
 R_{9}
 R_{10}
 R_{10}

55 (VI)

STEP C

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$$(V) + R_{11} - O - CO - O - CO - O - R_{11}$$
 R_{9}
 C
 CH_{2}
 R_{11}
 C
 CH_{2}

STEP D:

Compound V is allowed to react with an acyl chloride of the formula R_{II}-CO-Cl in a routine manner known to the art to afford a compound of Formula VIII.

$$(V) + R_{11} - CO - CI \longrightarrow X$$

$$R_{11}$$

$$(VIII)$$

STEP E:

A compound of Formula IX obtained from STEP B is subjected to a cleavage reaction to afford a compound of Formula X. Typically, to this end, compound IX is allowed to react with BBr₃/tetrahydroturan complex and the resultant product is hydrolyzed in a routine manner known to the art.

$$\begin{array}{c} \text{CH}_3\text{O} \\ \text{X} \\ \\ \text{R}_{10}\text{-H} \end{array} \begin{array}{c} \text{CH}_2\text{O} \\ \\ \text{R}_{10}\text{-H} \end{array}$$

(IX)

(X)

STEP F:

35 As a special case, a compound of Formula XI is allowed to react with chloroacetyl chloride in the presence of aluminum chloride in a routine manner known to the art to afford a compound of Formula XII (Friedel-Crafts reaction).

(XI)

STEP G:

Compound XII is allowed to react with a peracid, preferably m-chloroperbenzoic acid in a routine manner known to the art to afford a compound of Formula XIII (Baeyer-Villiger reaction).

(XIII)

STEP H:

Compound XIII is hydrolyzed preferably in the presence of a base such as sodium hydroxide to afford a compound of Formula XIV.

$$(XIII) + H_2O/NaOH$$

$$R_{10} \qquad 0$$

$$(XIV)$$

STEP I:

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A compound of Formula XV, where R_{1,8} is hydrogen, methoxy or hydroxy, which is obtained from one of the foregoing steps is allowed to react with hydroxylamine hydrochloride in a routine manner known to the art to afford a compound of Formula XVI. Typically, this reaction is conducted by first suspending compound XVI in othanol and thereafter adding an aqueous solution of sodium acotate and an aqueous solution of hydroxylamine hydrochloride to the suspension and stirring the resultant mixture at a temperature of 25 to 150°C.

STEP J

Compound XVI is allowed to react with an amino-C1-C6-alkyl bromide of the formula, Br-R13-NH2, where -R13-

(XVI)

NH₂ is an amino-C₁-C₆-alkyl group, in a routine manner known to the art to afford a compound of Formula XVII.

$$(XVI) + Br - R_{13} - NH_{2}$$

$$X$$

$$R_{3}$$

$$O - R_{13} - NH_{2}$$

$$(XVII)$$

STEP K

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Compound XV is allowed to react with a primary amine of the formula

where R₁₄ is C₁-C₅-alkyl, C₂-C₅-alkenyl, C₃-C₅-alkynyl, C₃-C₇-cycloalkyl, C₃-C₇-cycloalkyl, C₃-C₇-cycloalkyl or phenyl. Wherein the phenyl group may be substituted as indicated above, in a routine manner known to the art to afford an innine of Formula XVIII.

It is preferable to conduct this reaction in the presence of titanium (IV) isopropoxide and a suitable solvent such as acetonitrile. Typically, this reaction is conducted at a temperature of 0 to 80°C. This method is more advantageous than a method using TiCl₄ or a method wherein the reaction is conducted in a sealed tube at an elevated temperature with the aid of molecular sieves used as a water removing agent.

$$(XV) + H = \begin{pmatrix} R_{12} \\ R_{3} \end{pmatrix} \times \begin{pmatrix} CH_{2} \\ R_{3} \end{pmatrix}$$

STEP L:

Compound XVI is reduced with the aid of a Raney alloy and a sodium hydroxide solution in a similar manner as reported by B. Staskun and T. van Es (J. Chem. Soc., C., 531 (1966)) to afford a compound of Formula XIX.

STEP M:

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Compound XV is allowed to react with titanium isopropoxide and a secondary amine of the formula,

where the group

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NH NH
$$N-C_1-C_6$$
 -alky

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 — N N-phenyl or —N N-C₁-C₆-alkylphenyl

wherein the phenyl group may be substituted as indicated above.

followed by reduction with sodium cyanoborohydride under conditions similar to that described by R.J. Mattson et al., J. Org. Chem., 55, 2552-4 (1990), to afford a compound of Formula XX.

$$(XV) + H - NA$$

$$X$$

$$R_{12}$$

$$R_{3}$$

$$A$$

$$(XX)$$

STEP N

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Compound XVIII is reduced with sodium borohydride, sodium cyanoborohydride or borane/tetrahydrofuran complex in a routine manner known to the art to afford a compound of Formula XXI.

STEP O:

Compound XIX is reduced with the aid of borane/tetrahydrofuran and trifluoroacetic acid complex to afford a compound of Formula XXII.

$$(XIX) + BH_3/THF + CF_3CO_2H$$

$$X$$

$$X$$

$$R_3$$

$$NH_2$$

(XXII)

STEP P:

or 30

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A compound of Formula XXIII, which is obtained from STEP L or O, is allowed to react with a halide compound of the formula R₁₅-14 Mere P₁₅ is G-, C₂-alkyl, C₂-C₂-C₃-alkyl, C₁-C₂-c₃-alkyl, Phenyl-C₁-C₂-alkyl, wherein the phenyl group may be substituted as indicated above,

$$-Alk-N \qquad -Alk-N \qquad -Alk-N \qquad 0$$

-- Alk -- N --

to afford a compound of Formula XXIV.

R₁₂ (CH₂)_n + R₁₅ - Ha

(XXIII)

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$$\begin{array}{c|c} R_{12} & & & \\ & & & \\ X & & & \\ & & & \\ R_{3} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ & & \\ &$$

(XXIV)

STEP Q

A compound of Formula XXV, where R_{16} is hydrogen, C_3 – C_6 -allkyl, C_2 - C_6 -allknyl, C_3 - C_6 -allkyl, C_3 - C_7 -cycloalkyl, C_3 - C_7 -cycloalkyl, C_3 - C_7 -cycloalkyl, C_3 - C_7 -cycloalkyl, C_3 - C_7 -cycloalkyl, wherein the phenyl group may be substituted as indicated above, is allowed to react with formic acid in the presence of 1-(3-dimethylaminopyroyli-3-erbyl carbodilmide and 4-dimethylaminopyridine or the mixed anhydride prepared from formic acid and acetic anhydride to afford a compound of Formiula XXVII.

$$\begin{array}{c} R_{12} \\ X \\ R_{3} \\ R_{16} \end{array} CHO$$

STEP R

Compound XXV is allowed to react with an acyl chloride of the formula, R_{17} - CO - CI, where R_{17} is a C_1 - C_6 -alkyl group, in a routine manner known to the art to afford a compound of Formula XXVII.

$$(XXY) + R_{17} - CO - CT$$

$$X$$

$$R_{3}$$

$$R_{16}$$

$$C-C$$

$$R_{17}$$

$$(XXYII)$$

STEP S:

A compound of Formula XXV, where R₁₂ is not hydroxy, is allowed to react with a benzyl chloroformate in a routine manner known to the art to afford a compound of Formula XXVIII.

$$R_{12}$$

$$(CH_2)_n$$

$$R_3$$

$$R_{16}$$

$$C - C$$

$$CH_2$$

$$CH_2$$

$$CH_2$$

STEP T:

A compound of Formula XXV, where R_{12} is not hydroxy, is allowed to react with an isocyanate of the formula R_{17} . $\mathsf{N} = \mathsf{C} = \mathsf{O}$, where R_{17} is a $\mathsf{C}_1\text{-C}_{\mathsf{S}}$ -alloyt, phenyl- C_2 -charly group, wherein the phenyl group may be substituted as indicated above, to afford a compound of Formula XXIX. Typically, this reaction is conducted in the presence of a suitable catalyst such as 1,8-diazabicyoloj5.4.0|undeo-7-ene.

$$(XXV) + R_{17} - N - C - C$$

 $(R_{12} = -OH)$

25 STEP U:

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A compound of Formula XVI, where R_{12} is not hydroxy, is allowed to react with an isocyanate of the formula R_{17} - N=C=O in substantially the same manner as in STEP T to afford a compound of Formula XXX.

$$(XVI) + R_{17} - N = C - (R_{12} = -OH)$$

55 (XXX)

STEP V:

Compound XVI is allowed to react with an acyl chloride of the formula R₁₇ - CO - CI or an acid anhydride of the formula (R₁₇ - CO)₂O in a routine manner known to the art to afford a compound of Formula XXXI.

(XXXI)

STEP W:

A compound of Formula XXXII, where R_2 is not C_1 - C_6 -alkylarminocarbonyl, which is obtained from one of the foregoing STEPS is allowed to react with a chloroformate of the formula

in a routine manner known to the art to afford a compound of Formula XXXIII.

(XXXII)

$$\begin{array}{c} & & & \\ & &$$

40 STEP X:

A compound of Formula XXXIIIa which is obtained from STEP W is allowed to react with an isocyanate of the formula R₁₇-N₂-C₂O in substantially the same manner as in STEP T to afford a compound of Formula XXXIV Subsequently, Compound XXXIV is subjected to hydrogenolysis conducted with the aid of a suitable catalyst such as palladium-carbon in a routine manner known to the art to afford a compound of Formula XXXV.

(XXXIII)

$$\mathsf{C_6H_5CH_2} \bigvee_{\mathsf{O}}^{\mathsf{O}} \bigvee_{\mathsf{X}}^{\mathsf{O}} \bigvee_{\mathsf{R_3}}^{\mathsf{O}} \bigvee_{\mathsf{N}}^{\mathsf{CCH_2}})_{\mathsf{n}} \\ + \mathsf{R_17-N-C-O}$$

(XXXIIIa)

STEP Y:

A compound of Formula XXIII, which is obtained from one of the foregoing STEPS, is allowed to react with an isosyanate of the the formula $R_{17} \cdot N = C = O$ in substantially the same manner as in STEP T to afford a compound of Formula XXXVII.

(XXXXVI)

$$\begin{array}{c|c} & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & \\ & & & \\ & &$$

(XXXXII)

The compounds of Formula I and Formula III of the present invention are useful for the treatment of various memory dysfunctions characterized by a decreased cholinergic function such as Alzheimer's disease. Compounds of this invention also inhibit monoamine oxidase and/or act at central α_2 -adrenergic receptors and hence are useful as antidepressants.

The activity to alleviate such memory dysfunctions is manifested by the ability of these compounds to inhibit the enzyme acetylcholinesterase and thereby increase acetylcholine levels in the brain.

Cholinesterase Inhibition Assay

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Cholinesterases are found throughout the body, both in the brain and in serum. However, only brain acetylcholinesterase (AChE) distribution is correlated with central cholinergic innervation. This same innervation is suggested to be weakened in Alzheimer patients. We have determined in vitro inhibition of acetylcholinesterase activity in rat striatum.

In Vitro Inhibition of Acetylcholinesterase Activity in Rat Striatum

Acetylcholinesterase (AChE), which is sometimes called true or specific cholinesterase, is found in nerve cells, skeletal muscle, smooth muscle, various glands and red blood cells. AChE may be distinguished from other cholinesterases by substrate and inhibitor specificities and by regional distribution. Its distribution in brain roughly correlates with cholineratic innervation and subfractionation shows the highest level in nerve terminals.

It is generally accepted that the physiological role of AChE is the rapid hydrolysis and inactivation of acetylcholine.

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Inhibitors of AChE show marked cholinomimetic effects in cholinergically-innervated effector organs and have been used therapeutically in the treatment of glaucoma, myasthenia gravis and paralytic lieus. However, recent studies have succested that AChE inhibitors may also be beneficial in the treatment of Atzheimer's disease.

The method described below was used in this invention for assaying cholinesterase activity. This is a modification of the method of Ellman et al., Biochem, Pharmacol, 7, 88 (1961).

Procedure:

A. Reagents -

1. 0.05 M Phosphate buffer, pH 7.2

- (a) 6.85 a NaHoPOA·HoO/100 ml distilled HoO
- (b) 13.40 g Na₂HPO₄·7H₂O/100 ml distilled H₂O
 - (c) add (a) to (b) until pH reaches 7.2
 - (d) Dilute 1:10

2. Substrate in buffer

- (a) 198 mg acetylthiocholine chloride (10 mM)
- (b) q.s. to 100 ml with 0.05 M phosphate buffer,

pH 7.2 (reagent 1)

3. DTNB in buffer

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- (a) 19.8 mg 5,5-dithiobisnitrobenzoic acid (DTNB) (0.5 mM)
- (b) q.s. to 100 ml with 0.05 M phosphate buffer.

pH 7.2 (reagent 1)

4. A 2 mM stock solution of the test drug is made up in a suitable solvent and q.s. to volume with 0.5 mM DTNB (reagent 3). Drugs are serially diluted (1.10) such that the final concentration (in cuvette) is 10⁴M and screened for activity, If active, IC₅₀ values are determined from the inhibitory activity of subsequent concentrations.

B. <u>Tissue Preparation</u> -

Male Wistar rats are decapitated, brains rapidly removed, corpora striata dissected free, weighed and homogenized in 19 volumes (approximately 7 mg protein/ml) of 0.05 M phosphate buffer, pH 7.2 using a Potter-Elvehjem homogenizer. A 25 microliter aliquot of the homogenate is added to 1.0 milliter vehicle or various concentrations of the test drug and preincubated for 10 minutes at 37°C.

C. Assay -

Enzyme activity is measured with the Beckman DU-50 spectrophotometer. This method can be used for IC₅₀ determinations and for measuring kinetic constants.

Instrument Settings

Kinetics Soft-Pac Module #598273 (10) Program #6 Kindata:

50 Source - Vis

Wavelength - 412 nm

Sipper - none

Cuvettes - 2 ml cuvettes using auto 6-sampler

Blank - 1 for each substrate concentration

Interval time - 15 seconds (15 or 30 seconds for kinetics)

Total time - 5 minutes (5 or 10 minutes for kinetics)

Plot - yes

Span - autoscale

Slope - increasing Results - yes (gives slope)

Factor - 1

Reagents are added to the blank and sample cuvettes as follows:

Blank 0.8 ml Phosphate Buffer/DTNB

Control
 0.8 ml Phosphate Buffer/DTNB/Enzyme

0.8 ml Buffer/Substrate

0.8 ml Phosphate Buffer/Substrate

Drug: 0.8 ml Phosphate

15 Buffer/DTNB/Drug/Enzyme 0.8 ml Phosphate Buffer/Substrate

Blank values are determined for each run to control for non-enzymatic hydrolysis of substrate and these values are automatically subtracted by the kindata program available on kinetics soft-pac module. This program also calculates the rate of absorbance change for each curvette.

For IC₅₀ Determinations:

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Substrate concentration is 10 mM diluted 1:2 in assay yielding final concentration of 5 mM. DTNB concentration is 0.5 mM yielding 0.25 mM final concentration.

IC₅₀ values are calculated from log-probit an alysis.

30 Results of this assay for some of the compounds of this invention and physostigmine (reference compound) are presented in Table 1.

TABLE 1

35	Compound	Inhibitory Concentration, IC ₅₀ (μM) Brain AChE	
55	3-cyclopropylamino-4-methyl-1,2,3,4-tetrahydrocyclopent[b] indoi-7-yl methylcarbamate	3.5	
40	4-methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent[b] indoi-7-yl methylcarbamate	1.1	
	4-methyl-3-phenylmethylimino-1,2,3,4-tetrahydrocyclopent[b] indol-7-yl methylcarbamate	6.2	
45	Physostigmine	0.006	

This utility is further demonstrated by the ability of these compounds to restore cholinergically deficient memory in the Dark Avoidance Assay described below.

Dark Avoidance Assay

In this assay mice are tested for their ability to remember an unpleasant stimulus for a period of 24 hours. A mouse is placed in a chamfter that contains a dark compartment, it strong incandescent light drives it to the dark compartment, where an electric shock is administered through metal plates on the floor. The animal is removed from the testing apparatus and tested again, 24 hours later, for the ability to remember the electric shock.

If scopolamine, an anticholinergic that is known to cause memory impairment, is administered before an animal's initial exposure to the test chamber, the animal re-enters the dark compartment shortly after being placed in the test chamber 24 hours later. This effect of scopolamine is blocked by an active test compound, resulting in a greater interval before re-entry into the dark compartment.

The results for an active compound are expressed as the percent of a group of animals in which the effect of scopolamine is blocked, as manifested by an increased interval between being placed in the test chamber and reentering the dark compartment.

Results of this assay for some of the compounds of this invention and those for tacrine and pilocarpine (reference compounds) are presented in Table 2.

TABLE

		TABLE 2	
0	Compound	Dose (mg/kg of body weight, s.c)	% of Animals with Scopolamine Induced Memory Deficit Reversal
	3-cyclopropylamino-4-methyl-	0.63	27%
5	1,2,3.4-tetrahydrocyclopent[b]indol- 7-yl methylcarbamate	2.5	33%
	Tacrine	0.63	13%
0	Pilocarpine	5.0	13%

The utility is further demonstrated by the ability of these compounds to inhibit the enzyme monoamine oxidase, increase the brain levels of biogenic amine(s), and act as antidepressants.

Inhibition of Type A and Type B Monoamine Oxidase Activity in Rat Brain Synaptosomes

Purpose

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To determine the selective inhibition of the two forms of monoamine oxidase (MAO).

introduction

The metabolic dearnination of amines has been known for described two forms of monoamine oxidase, which are called "type A" and "type B". The existence of the two forms is based on different substrate and inhibitor specificities. Scrotonin (6HT) and norepinephrine (NE) are substrates for type A MAO, β-phenethylamine (PEA) and benzylamine are substrates for type B MAO, while dopamine (DA) and tyramine are substrates for both types. Clorgyline is a selective inhibitor of the type B enzyme and tranylocypromine and promized far en nonselective inhibitors of the type B enzyme and tranylocypromine and promized far en nonselective inhibitors.

Although various methods for measuring MAO activity are available, the described method involves the extraction of the radiolabeled dearminated metabolites of [91]—17 or [14C]—Phonenthylamine. This procedure allows MAO-A and MAO-B activities to be measured either simultaneously or individually (3).

Procedure

A. Reagents

1. Phosphate buffer (0.5 M), pH 7.4.

134.4 g NaH₂PO₄,7H₂O q.s. to 1 liter in distilled H₂O (A) 17.3 g Na₂HPO₄ q.s. to 250 ml in distilled H₂O (B) Adjust pH of A to 7.4 by slowly adding B (volumes as needed) Dilute 1:10 in distilled H₂O (0.05 M PO₄ buffer, pH 7.4)

- 2. 0.25 M Sucrose (PO₄ buffered):
 - 21.4 a sucrose, a.s. to 250 ml with 0.05 M PO₄ buffer
- 3. Substrate for MAO-A:
 - a. Serotonin creatine SO₄ (5HT) is obtained from Sigma Chemical Company. A 5 mM stock solution is made

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up in 0.01 N HCI. This is used to dilute the specific activity of the [3H]-5HT.

- b. [3H]-5-Hydroxytryotamine binoxalate (20-30 Ci/mmol) is obtained from New England Nuclear.
- c. Add 12 μ l of [3 H]-5HT to 2 ml of the 5 mM 5HT solution. (Final amine concentration in the assay is 200 μ M: see below.)

4 Substrate for MAO-B

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- a. β-phenethylamine (PEA) is obtained from Sigma Chemical Company. A 5 mM stock solution is made up in 0.01 N HCI. This is used to dilute the specific activity of the f¹⁴CI-PEA.
 - b B-[ethyl-1-14C]-phenethylamine hydrochloride (40-50 mCi/mmol) is obtained from New England Nuclear
 - c. Add 12 μ l of [¹⁴C]-PEA to 2 ml of the 5 mM PEA solution. (Final amine concentration in the assay is 200 μ M; see below.)
- 5. Equal amounts of MAO-A (5HT) and MAO-B (PEA) substrates are combined for simultaneously testing both MAO types, i.e. mixed stock solution of 2.5 mM 5HT and 2.5 mM PEA, 40 µl of this mixed solution gives a 200 µM final concentration of each amine in the assay. When testing only one MAO type, the individual 5 mM stock solutions must be diluted 1:1 with distilled water prior to adding 40 µl to the incubation mixture; i.e., same 200 µM final amine concentration.

20 B. Tissue Preparation

Male Wistar rats weighting 15:250 grams were seartficed and the brains rapidly removed. Whole brain minus cerebellum was homogenized in 30 volumes of ice-cold, phosphate-buffered 0.25 M sucrose, using a Potter-Elveim homogenizer. The homogenate was centrifuged at 1000 g for 10 minutes and the supernatant (S₁) decented and recentrifuged at 16,000 g for 20 minutes. The resulting pellet (P₂) was resuspended in fresh 0.25 M sucrose and served as the tissue source for mitochondrial MAO.

C. Assay

10 μl 0.5 M PO₄ buffer, pH 7.4

50 µl H₂O or appropriate drug concentration

400 µl Tissue suspension

Tubes are preincubated for 15 minutes at 37°C and the assay is started by adding 40 µl of combined substrate (1941bit1 and [140]-EA) at 15 second intervals. The tubes are incubated for 30 minutes at 37°C and the reaction stopped by the addition of 0.3 mi 2N HCI. Tissue blank values are determined by adding the scid before the radioactive substrate. The oxidative products of the reaction are extracted with ethyl acetate/folluene (1:1). 5 ml of this mixture is added to the tubes, The resultant mixture is vortexed for 15 seconds to extract the dearninated metabolites into the organic phase and the latter is allowed to separate from the aqueous phase. The tubes are placed in acetone/dry ice bath to freeze the acquous layer. When this layer is forzen, the top organic layer is poured into a scintillation value. 10 ml of Liquiscint is added and the samples are counted using window settings for ¹⁴C in one channel and ³H in the second channel. ICg₀ values are determined by log-probit analysis.

References

- 1. Johnston, J.P.: Some observations upon a new inhibitor of monoamine oxidase in brain tissue. Biochem. Pharmacol. 17: 1285-1297 (1968)
- Fowler, C. J. and Ross, S.B.: Selective inhibitors of monoamine oxidase A and B: biochemical, pharmacological and clinical properties. Med. Res. Rev. 4: 323-328 (1984).
- Kindt, M.V., Youngster, S.K., Sonsalla, P.K., Duvoisin, R.C. and Heikkila, R.E.: Role of monoamine oxidase-A (MAO-A) in the bioactivation and nigrostriatal dopaminergic neurotoxicity of the MPTP analog, 2'Me-MPTP. Eur. J. Pharmacol. 46: 313-318 (1988).

Results of the monoamine oxidase inhibition assay for representative compounds of this invention are presented in Table 3.

TABLE 3

	Compound	Inhibitory Concentration MAO-A	- IC ₅₀ (μM) MAO-B
5	1,2,3,4-Tetrahydrocyclopent[b]indol-3-(2-propynyl) amine	0 29	0,32
10	3-Cyclopropylamino-4-methyl- 1,2,3.4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate	0 32	0.42
15	4-Methyl-3-[(2-phenylcyclopropyl)imino]- 1,2,3.4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate	15.2	3.7
	(Reference Compounds)		
	Deprenyl	0.14	0.016
20	Tranylcypromine	0.19	0.12

The present inventors have also conducted Clonidine Binding Assay described below in order to ascertain the interaction of the compounds of this invention with α_2 -receptors.

³H-Clonidine Binding: α₂-Receptor

Introduction:

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A number of antidepressants have been shown to enhance neuronal release of norepinephrine by a presumed presynaptic α_2 -receptor blockade and this property may be of significance with respect to the mechanism of action of these compounds. See references 1, 2 and 3 cited below. The interaction of a compound with central α_2 -receptors is assessed in the ⁹t-clonidine binding assay.

35 <u>Procedure</u>

A. Reagents

1.

a. 57.2 g Tris HCI
16.2 g Tris Base - q.s. to 1 liter (0.5 M Tris buffer, pH 7.7)
b. Make a 1:10 dilution in distilled H₂O (0.05 M Tris buffer, pH 7.7)

45 2. Tris buffer containing physiological ions

a Stock Buffer

NaCl	7.014 g 0.372 g 0.222 g - q.s. to 100 ml in 0.5 M Tris buffer 0.204 g
KCI	0.372 g
CaCl ₂	0.222 g - q.s. to 100 ml in 0.5 M Tris buffer
MgCl ₂	0.204 g

- b. Dilute 1:10 in distilled H₂O. This yields 0.05 M Tris HCl, pH 7.7; containing NaCl (120 mM), KCl (5mM), CaCl₂ (2 mM) and MgCl₂ (1 mM)
- 3. [4-3H]-Clonidine hydrochloride (20-30 Ci/mmol) is obtained from New England Nuclear. For IC_{50}

determinations; 3H-Clonidine is made up to a concentration of 120 nM and 50 µl added to each tube (yields a final concentration of 3 nM in the 2 ml volume assay).

- 4. Clonidine-HCI is obtained from Boehringer Ingelheim. A stock solution of 0.1 mM clonidine is made up to determine nonspecific binding. This yields a final concentration of 1 µM in the assay (20 µl to 2 ml).
- 5. Test compounds. For most assays, a 1 mM stock solution is made up in a suitable solvent and serially diluted. such that the final concentration in the assay ranges from 10⁻⁵ to 10⁻⁸M. Seven concentrations are used for each assay and higher or lower concentrations may be used, depending on the potency of the drug.

B. Tissue Preparation

Male Wistar rats are sacrificed by decapitation and the cortical tissue rapidly dissected. The tissue is homogenized in 50 volumes of 0.05 M Tris buffer, pH 7.7 (buffer 1b) with the Brinkman Polytron, then centrifuged at 40,000 g for 15 minutes. The supernatant is discarded and the pellet re-homogenized in the original volume of 0.05 M Tris buffer. pH 7.7 and re-centrifuged as before. The supernate is discarded and the final pellet re-homogenized in 50 volumes of buffer 2b. This tissue suspension is then stored on ice. The final tissue concentration is 10 mg/ml. Specific binding is 1% of the total added ligand and 80% of total bound ligand.

C. Assay

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	100 µl	0.5 M Tris-physiological salts, pH 7.7 (buffer 2a)
	830 µl	H ₂ O
	20 µl	Vehicle (for total binding) or 0.1 mM clonidine (for nonspecific binding) or appropriate drug concentration
25	50 μl	³ H-clonidine stock
20	1000 µl	tissue suspension

Tissue homogenates are incubated for 20 minutes at 25°C with 3 nM 3H-clonidine and varying drug concentrations. and thereafter immediately filtered under reduced pressure on Whatman GF/B filters. The filters are washed with three five mI volumes of ice-cold 0.05 M Tris buffer, pH 7.7, and thereafter transferred to scintillation vials. Ten mI of Liquiscint counting solution is added to each sample which is then counted by liquid scintillation spectroscopy. Specific clonidine binding is defined as the difference between total bound and that performed using log-probit analysis. The percent inhibition at each drug concentration is the mean of triplicate determinations.

References

- 1. P.F. VonVoigtlander, "Antidepressant and Antipsychotic Agents", in "Annual Reports in Medicinal Chemistry", F. H. Clarke, ed., Chapter 1, Academic Press, New York, N.Y. (1976);
- 2. S. Clements Jewery, Neuropharmacol., 17, 779 (1978);
- 3. C.B. Smith and P. J. Hollingsworth, "Adrenergic Receptors and the Mechanism of Action of Antidepressant Treatments" in "Biochemical and Pharmacological Aspects of Depression", K.F. Tipton and M.B.H. Youdim, eds., Taylor and Francis, New York, N.Y., Chapter 4 (1989).

Results of the ³H-Clonidine Binding Assay for representative compounds of this invention are presented in Table 4.

TABLE 4

³ H-Clonidine Binding	
Compound	IC ₅₀ (μM)
1,2,3,3a,4,8a-Hexahydrocyclopent[b]indol-3-amine 2-naphthalenesulfonate hemihydrate	1.27
4-Methyl-1,2,3,4-tetrahydrocyclopent[b]indol-3-amine	1.49
4-Methyl-3-phenylmethylimino-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol	0.85
(Reference Compound)	
Amitriptyline	3.9

Effective quantities of the compounds of the invention may be administered to a patient by any of the various methods, for example, orally as in capsule or tablets, parenterally in the form of sterile solutions or suspensions, and

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in some cases intravenously in the form of sterile solutions. The free base final products, while effective themselves. may be formulated and administered in the form of their pharmaceutically acceptable acid addition salts for purposes of stability, convenience of crystallization, increased solubility and the like.

Acids useful for preparing the pharmaceutically acceptable acid addition salts of the invention include inorganic acids such as hydrochloric, hydrobromic, sulfuric, nitric, phosphoric and perchloric acids, as well as organic acids such as tartaric, citric, acetic, succinic, maleic, fumaric 2-naphthalenesulfonic and oxalic acids.

The active compounds of the present invention may be orally administered, for example, with an inert diluent or with an edible carrier, or they may be enclosed in gelatin capsules, or they may be compressed into tablets, For the ourgose of oral therapeutic administration, the active compounds of the invention may be incorporated with excipients and used in the form of tablets, troches, capsules, elixirs, suspensions, syrups, wafers, chewing gum and the like These preparations should contain at least 0.5% of active compounds, but may be varied depending upon the particular form and may conveniently be between 4% to about 70% of the weight of the unit. The amount of active compound in such compositions is such that a suitable dosage will be obtained. Preferred compositions and preparations according to the present invention are prepared so that an oral dosage unit form contains between 1.0 - 300 milligrams of active compound

The tablets, pills, capsules, troches and the like may also contain the following ingredients: a binder such as microcrystalline cellulose, gum tragacanth or gelatin; an excipient such as starch or lactose, a disintegrating agent such as alginic acid, Primogel, comstarch and the like; a lubricant such as magnesium stearate or Sterotex; a glidant such as colloidal silicon dioxide; and a sweeting agent such as sucrose or saccharin may be added or a flavoring agent such as peppermint, methyl salicylate, or orange flavoring. When the dosage unit form is a capsule, it may contain, in addition to materials of the above type, a liquid carrier such as a fatty oil. Other dosage unit forms may contain other various materials which modify the physical form of the dosage unit, for example, as coatings, Thus, tablets or pills may be coated with sugar, shellac, or other enteric coating agents. A syrup may contain, in addition to the active compounds, sucrose as a sweetening agent and certain preservatives, dyes, coloring and flavors. Materials used in preparing these various compositions should be pharmaceutically pure and non-toxic in the amounts used.

For the purpose of parenteral therapeutic administration, the active compounds of the invention may be incorporated into a solution or suspension. These preparations should contain at least 0.1% of active compound, but may be varied between 0.5 and about 30% of the weight thereof. The amount of active compound in such compositions is such that a suitable dosage will be obtained. Preferred compositions and preparations according to the present inventions are prepared so that a parenteral dosage unit contains between 0.5 to 100 milligrams of active compound.

The solutions or suspensions may also include the following components; a sterile diluent such as water for injection, saline solution, fixed oils, polyethylene glycols, glycerine, propylene glycol or other synthetic solvents; antibacterial agents such as benzyl alcohol or methyl parabens; antioxidants such as ascorbic acid or sodium bisulfite; chelating agents such as ethylenediaminetetraacetic acid; buffers such as acetates, citrates or phosphates and agents for the adjustment of tonicity such as sodium chloride or dextrose. The parenteral preparation can be enclosed in disposable syringes or multiple dose vials made of glass or plastic.

Examples of the compounds of this invention include:

1,2,3,4-tetrahydrocyclopent[b]indol-3-amine;

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- 4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-3-amine;
 - 1,2,3,4-tetrahydrocyclopent[b]indol-3-cyclopropylamine;
 - 4-methyl-1,2,3,4-tetrahydrocyclopentfblindol-3-cyclopropylamine
 - 1.2.3.4-tetrahydrocyclopent(blindol-3-(2-propynyl)amine:
 - 1,2,3,4-tetrahydrocyclopent[b]indol-3-(N-formyl)amine;
 - 1.2.3.4-tetrahydrocyclopent(blindol-3-(N-phenylmethyloxycarbonyl)amine;
 - 1,2,3,3a,4,8b-hexahydrocyclopent[b]indol-3-amine;
 - 1.2.3.3a.4.8b-hexahvdro-4-methylcyclopent[b]indol-3-amine:
 - 1,2,3,3a,4,8b-hexahydro-4-methylcyclopent[b]indol-3-(2-propynyl
 - 4-methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent/blindo
 - 4-methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent[b]indo 7-vl methylcarbamate;
 - 3-(N-cyclopropyl)amino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol;
 - 3-(N-cyclopropyl)amino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate;

 - 3-cyclopropylamino-1,2,3,3a,4,8b-hexahydro-4-methylcyclopent[b]indol-7-ol; 3-cyclopropylamino-1,2,3,3a,4,8b-hexahydrocyclopent[b]indol-7-vl phenylmethylcarbonate:
 - 3-(N-cyclopropyl-N-methylaminocarbonyl)amino-1,2,3,3a,4,8b-hexahydrocyclopent[b]indol-7-yl phenylmethylcar-
 - 3-(N-cyclopropyl-N-methylaminocarbonyl)amino-1,2,3,
 - 3a,4,8b-hexahydro-4-methylcyclopent[b]indol-7-ol;

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3-cyclopropylamino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate; 1.2.3.3a.4.8b-hexahvdro-4-methyl-3-phenylmethylaminocyclopent(blindol-7-ol: 1.2.3.3a.4.8b-hexahydro-4-methyl-3-aminocyclopent(blindol-7-ol: 1,2,3,3a,4,8b-hexahydro-4-methyl-3-phenylmethyloxycarbonylaminocyclopent[b]indol-7-ol; 1.2.3.3a.4.8b-hexahvdro-4-methyl-3-(N-phenylmethyloxycarbonyl)amino-cyclopent(blindol-7-yl methylcarbamate) 1,2,3,3a,4,8b-hexahydro-4-methyl-3-methylaminocarbonylaminocyclopent[b]indol-7-ol; 1,2,3,3a,4,8b-hexahydro-4-methyl-3-(N-phenylmethyl-N-methylaminocarbonyl)aminocyclopent[b]indol-7-ol; 4-t-butyloxycarbonyl-1.4-dihydrocyclopent(b)indol-3(2H)-one; 7-chloroacetyl-1,4-dihydro-4-methylcyclopent[b]indol-3(2H)-one; 7-chloroacetyloxy-1 4-dihydro-4-methylcyclopent[b]indol-3(2H)-one; 1,4-dihydro-7-hydroxy-4-methylcyclopent(b]indol-3(2H)-one; 1.4-dihydro-7-methylaminocarbonyloxy-4-methylcyclopentfblindol-3(2H)-one: 3-hydroxylimino-7-methoxy-1.2.3.4-tetrahydrocyclopent[b]indole; 3-hydroxylimino-1,2,3,4-tetrahydrocyclopent[b]indole; 3-hydroxylimino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indole; 3-(2-aminoethyl)oximino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indole; 3-cvclopropylimino-1,2,3,4-tetrahydrocyclopent[b]indole: 3-cyclopropylimino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol

3-hydroxylimino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-3-acetyloxylimino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl-acetate; 4-methyl-3-penyloxthylimino-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl-acetate;

4-methyl-3-phenylmethylimino-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol

4-methyl-3-phenylmethylimino-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate;

4-methyl-3-[(2-phenylcyclopropyl)imino]-1,2,3,4-tetrahydrocyclo [b]indol-7-ol;

4-methyl-3-[(2-phenylcyclopropyl)imino]-1,2,3,4-tetrahydrocyclo [b]indol-7-yl methylcarbamate; 3-cyclopropylimino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol 7-ol;

3-methylaminocarbonyloximino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate;

3-cyclopropylamino-1,2,3,3a,4,8b-hexahydro-4-methylcyclopent[b] indol-7-yl methylcarbamate; 3-amino-1,2,3,3a,4,8b-hexahydro-4-methylcyclopent[b]indol-7-yl 1,2,3,4-tetrahydroisoquinolylcarbamate;

5-bromo-3-cyclopentylamino-1,2,3,3a,4,8a-hexahydro-4-methylcyclopent[b]indol-7-yl phenylethylcarbamate, 3-[2-morpholinoethylamino]-4-methyl-1,2,3,4-tetahydrocyclopent indol-7-yl phenylethylcarbamate; and

4-methyl-3-(4-piperdinyl)amino-1,2,3,4-tetrahydrocyclopent/blindol-7-yl phenylethylcarbamate.

EXAMPLE 1

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3-Hydroxyimino-7-methoxy-1,2,3,4-tetrahydrocyclopent[b]indole

A stirred solution of 1,2-cyclopentadione mono-4-methoxypheny/hydrazone (6.0 g) in 100 ml of 10% aqueous H₂SQ₄ was heated on a steam bath for 4 hours and thereafter allowed to cool to room temperature and filtered to give 1,4-dihydro 7-methoxy-cyclopentlibjindoi-(2(1)-no ea as solid. To the indole (2.6 g) in 25 ml of 95% E10H was added hydroxylamine hydrochloride (1.7 g) in 15 ml water followed by sodium acetate (2.1 g) in 25 ml of 95% e10H was ended at reflux for 2.5 hours and allowed to stand overnight. The E10H was removed in 15 ml vaccor and the solid material which formed was collected and purified by flash chromatography to give 0.8 g of a mixture of oxime isomers.

ANALYSIS:			
Calculated for C ₁₂ H ₁₂ N ₂ O ₂	66.65%C	5.59%H	12.95%N
Found	66.39%C	5.51%H	12.91%N

EXAMPLE 2

3-Hydroxyimino-1,2,3,4-tetrahydrocyclopent[b]indole

To a stirred solution of 1.4-dihydrocyclopentiblindol-3(2H)-one* (10 g) in 100 mt of 95% E1OH was added hydrocylyamine hydrochioride (8.3 g) in 20 ml water followed by sodium acetate (9.7 g) in 20 ml water. The mixture was heated at reflux for 2 hours and thereafter allowed to stand at room temperature overnight. The E1OH was removed in vacuo and the solid material which formed was collected and recrystallized from 95% E1OH to give 4.5 g of predominantly one oxime isomer in the first crop and 3.0 g of a mixture of oxime isomers from the second crop. A 1.5 g sample of the *Elios et al., J. Chem. Soc., 634 (1944). single isomer was recrystallized to provide 0.9 g of analytically pure material.

ANALYSIS:			
Calculated for C ₁₁ H ₁₀ N ₂ O	70.95%C	5.41%H	15.04%N
Found	70.71%C	5.32%H	14.94%N

EXAMPLE 3

3-Hydroxyimino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indole

To a stirred solution of 1,4-dihydro-4-methylcyclopent[b]indol-3(2H)-one (3.0 g) in 30 ml of 95% EIOH was added hydroxylamine hydrochloride (2.25 g) in 9 ml water followed by sodium acetate (4.4 g) in 9 ml water. The mixture was heated at reflux for 4 hours and threatler an additional 1.1 gram of hydroxylamine hydrochloride in 5 ml water and 2.2 grams of sodium acetate in 5 ml water were added. After an additional 2 hours of reflux, the mixture was allowed to stand at room temperature overnight. The material which precipitated was collected and recrystallized from 95% EIOH to give 1.9 grams of analytically pure material.

ANALYSIS:			
Calculated for C ₁₂ H ₁₂ N ₂ O	77.98%C	6.04%H	13.99%N
Found	72.18%C	6.11%H	14.00%N

EXAMPLE 4

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3-(2-Aminoethyl)oximino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indole

To a stirred suspension of 3-hydroxyimino-4-methyll 2,3 41-terlahydroxyclopent(blindole (5.0 g) in methylene chloride (50 ml) was added 50% NaOH (50 ml) followed by tetrabutylammonium bromide (800 mg) and bromoethylamine hydrobromide (7.6 g). The reaction mixture was stirred overnight at room temperature. The layers were separated and the aqueous layer was extracted with methylene chloride (50 ml). The organic layers were combined, dried (Na₂SO₄) and concentrated. The product was purified via flash chromatography on silica gel eluting with 10% methanol/methylene chloride to provide 1.1 grams of purified material.

25 EXAMPLE 5

1,2,3,4-Tetrahydrocyclopent[b]indol-3-amine

To a stirred solution of 3-hydroxyrimino-1,2,3,4-tetrahydroxyclopentlylindole (6 g) in 150 ml of 95% EIOH at 0°C was added a nickel alloy (Harehaw, Ni-1000P, 10 g) followed by 12.9 grams of sodium hydroxide in 150 ml water. The ice bath was removed after 0.5 hour and the mixture was stirred an additional hour and filterof. The EIOH was removed in vacuo and the product crystallized to provide 5.0 grams of solid. A sample was recrystallized from toluene to provide analytically pure material.

L	ANALYSIS:			
Γ	Calculated for C ₁₁ H ₁₂ N ₂	76.71%C	7.02%H	16.26%N
ı	Found	76.44%C	6.98%H	15.99%N

50 EXAMPLE 6

4-Methyl-1,2,3,4-tetrahydrocyclopent[b]indol-3-amine hydrochloride

To a stirred solution of 3-hydroxylimino-4-methyl-1,2,34-tetrahydroxyleopent[b]indole (\$ g) in 200 ml 95% EICH at 0°C was added a nickel alloy (9 g) followed by 11 grams of sodium hydroxide in 200 ml water. The ice bath was removed after 0.25 hours and the mixture was stirred an additional hour. Additional nickel alloy (2x 1 gram) was added and the mixture was stirred for 2 hours. The catalyst was removed by filtration, the EICH was removed in vacuo and the product extracted into CHyE₂Ck (2x 100 ml). The CH₂Ck (2x 100 ml).

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The oil (2.0 g) was dissolved in diethyl ether (100 ml) and ethereal HCl was added until the solution became slightly acidic. The solid which formed was filtered and dried overnight to provide 1.6 grams of 4-methyl-1,2.3,4-tetrahydrocy-closentiblindoi-3-amine hydrochloride.

ANALYSIS:				
Calculated for C ₁₂ H ₁₄ N ₂ ·HCI 64.72%C 6.79%H 12.58%N				
Found	64.41%C	6.82%H	12.18%N	

EXAMPLE 7

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4-t-Butyloxycarbonyl-1,4-dihydrocyclopent[b]indol-3(2H)-one

To a stirred solution of 1,4-dihydrocyclopent[b]indol-3(2H)-one (10.0 g) in acetonitrile (100 ml) was added diabutylpyrocarbonate (15 g), followed by 4-dimethylaminopyridine (700 mg). The mixture was stirred overnight at room temperature under nitrogen. The solvent was removed under reduced pressure and the residue was purified by flash column chromatography to provide 4-t-butyloxycarbonyl-1,4-dihydrocyclopent[b]indole-3(2H)-one (4.5 g) as a solid.

ANALYSIS:			
Calculated for C ₁₆ H ₁₇ NO ₃	70.83%N	6.32%H	5.16%N
Found	71.04%C	6.35%H	5.16%N

EXAMPLE 8

1,2,3,4-tetrahydrocyclopent[b]indol-3-cyclopropylamine hydrochloride

1.4-dhydrocyclopen(Ib)indol-3(2H)-one (5 0 g) was separated into two portions and placed into sealed thube sech containing to lunes (20 ml), cylopropylamine (20 ml), and 25 ml policular sieves (1 g). The mixtures were placed in an oil bath and refluxed for 7 hours. Each tube was allowed to cool to ambient temperature, the molecular sieves were filtered, and the filtrate concentrated to give a brown solid which was identified as the imine via NMR/MS. The combined imine product was dissolved in isopropanol (125 ml) and methanol (25 ml), and thereafter sodium borohydride (2.66 g) was added and the mixture was stirated under nitrogen at ambient temperature overnight. The mixture was cooled to °C, water was slowly added and the mixture was stirated on brough. The stock (2x 200 ml), the ElOAc layer was extracted with 10% HCl (2x 200 ml) and the acid extracts were neutralized (10% NaCH) and concentrated in viacure to give 3.5 grams of product. A 1.5 gram sample was dissolved in EloC (100 ml) and ethereal HCl was added, the pracipitate was collected and dried to provide 1.23.4 a-tetratedyrocyclopentibilited-3-evelopropylamine hydrochloride.

ANALYSIS:			
Calculated for C ₁₄ H ₁₆ N ₂ ·HCI	67.60%C	6.89%H	11.26%N
Found	67.22%C	6.87%H	10.79%N

EXAMPLE 9

4-Methyl-1,2,3,4-tetrahydrocyclopent[b]indol-3-cyclopropylamine 2-naphthalene sulfonate

1.4-Dhydro-4-methyl-cyclopent[bi]ndol-3(2H)-one (2.0 g) and cyclopropylamine (3.0 g) were dissolved in 30 ml toluene and cooled to -10°C. Tilanium tetrachloride (0.7 m ill) was dissolved in 10 ml toluene and added to the first solution. The reaction mixture was allowed to warm to room temperature and stirred overnight. The imine was isolated by filtering the mixture through a paid of slica and removing the solvent in vacuor. The imine (2.4 g) was dissolved in 00 ml of 5:1 #PCH/MoCH and thereafter sodium borehydride (1.2 g) was added. The reaction mixture was stirred overnight. The solvents were removed and the product purified by chromatography isolating the product as a yellow oil (1.6 g).

A 0.75 g portion of the cyclopropylaminoindole compound was dissolved in 75 ml Et₆O and stirred while a solution of 0.69 g of 2-naphthalene sulfonic acid in 50 ml Et₆O was added slowly. A white precipitate formed which was filtered under N₂₀, washed with 2x 50 ml Et₆O and dried to afford 1.04 g of 4-methyl-1,2,34-tetrahydrocyclopen(filt)ndol-3-cy-

clopropylamine 2-naphthalene sulfonate.

ANALYSIS:			
Calculated for C ₁₅ H ₁₈ N ₂ ·C ₁₀ H ₈ SO ₃	69.10%C	6.03%H	6.45%N
Found	68.98%C	6.04%H	6.39%N

EXAMPLE 10

1.2.3.4-Tetrahydrocyclopent(blindol-3-(2-propynyl)amine

To a stirred solution of 1,2,3.4-tetrahydrocyclopent[bjindol-3-amine (5,0 g) in tetrahydrofuran (30 ml) under nitrogen was added triethylamine (2.9 g) followed by a dropwise addition of propargyl bromide (4.45 g, 60% solution in toulous) dissolved in tetrahydrofuran (20 ml). The mixture was stirred overnight. Additional propargyl bromide (0.01 mole) dissolved in tetrahydrofuran (10 ml) was added and the mixture was stirred for 3 hours. The mixture was concentrated in zezu, O,H-(2, (15 ml)) was added and the mixture was storelated with 10% Hol (12 x5 oml). The organic phase was dried (Na₂SO₄) and concentrated to give 0.85 gram of product. The reaction was repeated on the same scale using identical conditions. The products were combined and chromatographed on silica gel eluting with 5% MeOH/OH₂Cl₂ to provide 1.23, 4-tetrahydrocyclopent[bjindol-2(2-propyym)]amine (1.6 g).

ANALYSIS:			
Calculated for C ₁₄ H ₁₄ N ₂	79.97%C	6.71%H	13.32%N
Found	79.70%C	6.77%H	13.14%N

EXAMPLE 11

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1,2,3,4-Tetrahydrocyclopent[b]indol-3-(N-formyl)amine

To a stirred solution of 1,2,3,4-tetrahydrocyclopen(b)indol-3-amine (2.0 g) in 25 mt methylene chloride at room temperature was added 4-drinethylaminopriorilino (1.4 g) followed by 0.46 mt of formic acid. 1-(3-Dimethylaminopropyi)-3-ethylcarbodimide bydrochloride (2.4 g) was added and the mixture was stirred overnight under a nitrogen atmosphere. The reaction mixture was diluted with CH_2C_{12} (100 ml), extracted with water (3x 50 ml), dried (Na₂SO₄) and concentrated in vacuo to give a solid which was crystallized from EICH and recrystallized from toluene to provide 1.2.3.4-tetrahydrocyclopen(b)indol-3-(N-tomylamine (1.1 q.)

ANALYSIS:			
Calculated for C ₁₂ H ₁₂ N ₂ O	71.98%C	6.04%H	13.99%N
Found	71.91%C	5.86%H	13.54%N

EXAMPLE 12

1,2,3,4-Tetrahydrocyclopent[b]indol-3-(N-phenylmethyloxycarbonyl)amine

To a stirred solution of 1,2,3.4-tetrahydrocyclopent(b)indol-3-eninio (5 g) in 50 ml CH₂Cl₂ at room temperature was added trietly lamine (3.2 g) followed by 5.4 grams of benzyl chloroformate in 25 ml CH₂Cl₂. The mixture was stirred for 2 hours and thereafter washed successively with water (50 ml), 10% HCl (50 ml) and water (50 ml). The CH₂Cl₂ solution was dried (Na₂SO₄). Concentrated in vacuo and purified by flash chromatography eluting with 2:1 hexane/ action to give 2.0 grams 1,2,4-tetrahydrocylopent(b)indol-4.(N-penylmethydroxycarbonylparinie,

ANALYSIS:			
Calculated for C ₁₉ H ₁₈ N ₂ O ₂	74.49%C	5.92%H	9.14%N
Found	74.23%C	5.99%H	8.96%N

EXAMPLE 13

1,2,3,3a,4,8b-Hexahydrocyclopent[b]indol-3-amine 2-naphthalenesulfonate hemihydrate

1.2.3.4-Tetrahydrocyclopen(Ip)Indol-3-amine (2.0.9) was piaced in a three-neck flask under nitrogen and 34 ml of a 1.0 M borane-tetrahydrotural mass added dropwise via a syringe. The misture was stirred at 0°C for 0.5 hour and thereafter trifluroacetic acid (34 ml) was added in a dropwise manner. After stirring for 2 hours, the tetrahydrofuran was removed *In vacuo*, and the residue was made basic with 10% NaOH, extracted with CH₂OI₂ (2x 75 ml) and concentrated to an oil (2 grams). A 1.0 grams amplie of the oil was dissolved in either (200 ml) as solution of 1.3 grams of 2-napthalene sulfonic acid in other was added in a dropwise manner with stirring. The precipitate which formed was collected by filtration under nitrogen.

ANALYSIS:			
Calculated for C ₁₁ H ₁₅ N ₂ ·C ₁₀ H ₈ SO ₃ ·0.5H ₂ O	64.41%C	5.93%H	7.15%N
Found	64.34%C	5.33%H	6.73%N

EXAMPLE 14

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1,2,3,3a,4,8b-Hexahydro-4-methylcyclopent[b]indol-3-amine 2-naphthalenesulfonate

4-Methyl-1, 2, 3,4-tetrahydrocyclopent[b]indol-3-amine (10.2 g) was placed in a three-neck flask under nitrogen and 17 of a 1.0 M borane-letrahydrofuran complex in tetrahydrofuran was added dropwise via a syrings. The mixture was stirred at CPC for 0.5 hours and thereafter trifluorocectic acid (185 mi) was added via a pressure-addition fund. Alter stirring for 1 hour, the tetrahydrofuran was removed in vacuo, and the residue was basified with 10% NaOH (pH=8), extracted with 0H₂Cl₂ (2x 500 mi), dried over Na₂SO₄ and concentrated to an oil (10.3 g). The crude material was purified by column chromatography.

A 1.7 g sample of 1,2,3,3a,4,8b-hexahydro-4-methylcyclopent[b]indol-3-amine was dissolved in 150 ml Et₆O and a solution of 1.9 g of 2-naphthalene sulfonic acid in ether was added in a dropwise manner with stirring. A solid was collected by fittration under N₅

ANALYSIS:			
Calculated for C ₁₂ H ₁₇ N ₂ ·C ₁₀ H ₈ SO ₃	66.64%C	6.10%H	7.06%N
Found	66.74%C	5.66%H	6.77%N

EXAMPLE 15

1,2,3,3a,4.8b-hexahydro-4-methylcyclopent[b]indol-3-(2-propynyl)amine hydrochloride

1,2,3,3a,4,8b-Hexahydro-4-methylcyclopent[b]indol-3-amine (5.0 grams) was dissolved in 50 ml tetrahydrofuran aladiom (2.7 grams). The solution was cooled to 0°C and propagyl bromide (3.2 grams) in 20 ml tetrahydrofuran was added slowly. After the addition, the mixture was allowed to come up to room temperature and stirred overnight. The tetrahydrofuran was stripped off and the residue taken up in 200 ml CH₂Cl₂. The organic layer was extracted with 10% hG1 (2x 70 ml). The aqueous layer was extracted with 10% hG1 (2x 70 ml). The aqueous layer was extracted with 2x 200 ml CH₂Cl₂ and the organic layers were combined and dried over sodium sulfate. The solvent was removed <u>in vacuo</u>. Flash chromatography on silica gel gave 1,2,3,3a,4,8b-hoxshydro-4-methylcyclopent[b]indol-3-(2-propynylamine (2.0 grams) as a reddish brown oil.

A 1.46 g sample of the indoline was dissolved in ether and stirred vigorously. An ethereal HCl solution was added to this solution until neutral (pH=6). The solids were then filtered and dried under N₂ giving 1,2,3,3a,4,8b-hexahydro-4-methylcyclopent([b]ndol-2-(2-propynylpamine hydrochlonide as a fine white powder (1.46 g grams).

ANALYSIS:

Calculated for C₁₅H₁₈N₂*HCl: 68.56*C 7.30*H 10.68*N Found: 68.21*C 7.27*H 10.54*N

EXAMPLE 16

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7-Chloroacetyl-1,4-dihydro-4-methylcyclopent[b]indol-3(2H)-one

Aluminum chloride (6.5g) was suspended in $\mathrm{CH}_2\mathrm{Cl}_2$ (20 ml) at 0°C, chloroacetyl chloride (7.2 g) was slowly added and mixture was intered for 5 minutes. This mixture was added dropwise to a stirred solution of 1,4-dilydo-4-methyl-cyclopent[b]indoi-3(2H)-one (6.0 g) in 100 ml CH_2Cl_3 at 0°C. The mixture was stirred at 0°C for 45 minutes and thereafter an additional equivalent of preformed solution of aluminum chloride and chloroacetyl chloride in methylene chloride was introduced in a dropwise manner. After 30 minutes the reaction mixture was slowly poured into a stirred i cewater mixture. The layers were separated and the CH_2Cl_2 layer was washed with NaH-CO₃ dried (Na₂SO₄) and concentrated to an oil. Purification by flash chromatography on slica gel eluting with hexane/acetone provided 7-chloroacetyl-1,4-di-hydro-4-methyl-voclopent[b]indoi-3(2H)-one (4.5 g).

ANALYSIS:			
Calculated for C ₁₄ H ₁₂ CINO ₂	64.25%C	4.62%H	5.35%N
Found	64.35%C	4.61%H	5.24%N

25 EXAMPLE 17

7-Chloroacetyloxy-1,4-dihydro-4-methylcyclopent[b]indol-3(2H)-one

To a stirred solution of 7-chloroacetyl-1,4-dilhydro-4-methylcyclopent[b]indol-3(2H)-one (2.0 g) in chloroform (100 ml) was added sodium phosphate (1.0 2 g) followed by m-chloroperbenzole acid (2.5 g, 50-66% purity). The mixture was stirred at room temperature under a nitrogen atmosphere for 14 hours. Saturated NaHCO₃ aqueous solution (50 ml) was added, the layers separated and the organic layer washed with water (2x 50 ml). The solution was dried (Na₂SO₄), filtered and concentrated to give a yellow oil which crystallized upon standing. Recrystallization from with EIOH provided 7-chloroacetyloxy-1,4-dilhydro-4-methylcyclopyth[bidol-2(H)-one (1.1 q.)

ANALYSIS:

Calculated for C14H12ClNO3: 60.55%C 4.36%H 5.04%N

Found: 60.47%C 4.33%H 4.98%N

45 EXAMPLE 18

1,4-Dihydro-7-methylaminocarbonyloxy-4-methylcyclopent[b]indol-3(2H)-one

7-Chloroacelylosy-1,4-dilydro-4-methylcyclopentl[b]indol-3(2H)-one (5.0 g) was suspended in EICH (100 ml), and thereafter 10% NaCH solution (50 ml) was added and the mixture was stirred at room temperature for 3 hours. The mixture was concentrated in vacuo, CH₂Cl₃, (100 ml) was added followed by 10% HCl until the aqueous layer was neutralized The layers were separated and the aqueous phase extracted with CH₂Cl₂ (2x 100 ml). The organic portion was dired (Na₂SCl₄) and concentrated and the residue was recrystallized from 95% EICH to provide 1,4-dihydro-7-hydroxy-4-methylcyclopent[b]indol-3(2H)-one as an off-white solid. The phenol was disested in CH₂Cl₂ (100 ml), and thereafter 1,6-diszabicycl(5,4-d) quadec-7-ene (4,9) was added followed by methyl isocyvanet (1,4-g) and the mixture was stirred overnight. The mixture was stored overnight. The mixture was ocncentrated in vacuo to afford an oily solid which was crystallized from EICH to provide 1,4-dihydro-7-methylcynoy-4-methylcytopentib|Indol-3(2H)-one (1,1 a)

ANALYSIS:			
Calculated for C ₁₄ H ₁₄ N ₂ O ₃	65.11%C	5.46%H	10.85%N
Found	65.20%C	5.32%H	10.74%N

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3-Acetyloxyimino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl acetate

7-Chloroacelyloxy-1.4-dihydro-4-methylcyclopent(b)indoi-3(2H)-one (6.0 g) was suspended in EiOH (200 ml) and a solution of NaOAc (15.6 g) in water (25 ml) and a solution of hydroxylamine hydrochloride (6.0 g) in water (25 ml) were added and the mixture was refluxed for 3 hours. The fixture was concentrated in vazuo and the residue was recrystallized from 95% EiOH to provide 3-hydroxylimino-4-methyl-1.2,3.4-tetrahydrocyolopent(b)indoi-7-ol as an off-white solid 1 hot sonime was dissolved in tetrahydrotran (100 ml), and thereafter acetic anhydride (6.1 g) and 4-direkylaminopyridine (400 mg) were added and the mixture was stirred under nitrogen at ambient temperature overnight. The mixture was concentrated in vazuo, CH₂Cl₂ (100 ml) was added and the solution was washed successively with water (50 ml), 5% NaHCO₃ (50 ml) and water (50 ml). After drying (Na₂SO₄), the solvent was removed in vazuo and the product recrystallized from EiOH to provide 3-acetyloximino-4-methyl-1,2,3.4-tetrahydrocyclopen(b)indoi-7-yl-acetate (1.7 q).

ANALYSIS:			
Calculated for C ₁₆ H ₁₆ N ₂ O ₂	63.99%C	5.37%H	9.33%N
Found	63.56%C	5.37%H	9.29%N

EXAMPLE 20

4-Methyl-3-phenylmethylimino-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol

To a stirred suspension of 7-chloroscetyloxy-1.4-dihydro-4-methyloyolopent[b]-indol-3(2H)-one (6.0 g) in toluene (50 mi) was acided benzylamine (9.2 g) and the mixture was heated at reflux temperature with azeotropic removal of water using a Dean Stark trap. After 4 hours, TLC analysis indicated complete conversion to product. The mixture was allowed to cool to room temperature and filtered, and the solid material was washed with acetonlitrile. The filtrate and washings were combined, concentrated and purified by flash chromatography on silica gel (2.1 hexane/acetone as eluent). The crystals which formed in the product-containing fractions were collected via filtration to give 4-methyl-3-pheny/methyllimino-1.2,9.4-tetrahydrocyclopent[b]indol-7-ol (1.1 grams) and the filtrate was concentrated to give an oil (3.0 arrans) which crystallized upon standing.

ANALYSIS:			
Calculated for C ₁₉ H ₁₈ N ₂ O	78.59%C	6.25%H	9.65%N
Found	78.62%C	6.21%H	9.63%N

EXAMPLE 21

4-Methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol

To a stirred solution prepared from 4-methyl-5-phenylmethyllmino-1.2.3.4-tetrahydrocyclopent[b]indol-7-of (16.0 g), isopropanol (200 ml) and methanol (50 ml) was added sodium borohydride (4.8 g) and the mixture was stirred under nitrogen at ambient temperature for 3 hours. The mixture was cooled to 0°C, water was slowly added and the mixture was stirred 0.5 hour. The mixture was extracted with CH-5Cl₂ (2x.200 ml), and the CH₂Cl₂ extracts were dired (Na₂SO₄), concentrated and chromatographed on silica gel eluting with 2.1 hexanes/actions. The product-containing fractions were combined to give 4.25 grams of 4-methyl-3-(phenylmethylamino-) 1.2.3.4-tetrahydrocyclopent[b]indol-7-of.

ANALYSIS:			
Calculated for C ₁₉ H ₂₀ N ₂ O	78.05%C	6.89%H	9.58%N

(continued)

ANALYSIS:			
Found	78.20%C	6.97%H	9.54%N

EXAMPLE 22

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4-Methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate

To a strired solution of 4-methyl-5-phenylmethylimino-1.2.34-tetrahydrocyclopent[b]indoh7-ol (2.0 g) in CH₂Cl₂ (40 mi) was added 1,8-diazabicyclof,5.4.0[undec-7-ene (0.16 g) followed by the dropwise addition of methyl loscyanate (0.39 g) in CH₂Cl₂ (10 mi). The reaction was monitiored via TLC and after 3 hours the solution was concentrated and the precipitate was collected and recrystallized from acetonitrile to give 4-methyl-3-phenylmethylimino-1,2.3.4-tetrahydrocyclopent[bindot-7-q interhylcarbarent cl.1.85 crams).

ANALYSIS:	-		
Calculated for C ₂₁ H ₂₁ N ₃ O ₂	72.60%C	6.09%H	12.09%N
Found	72.59%C	6.01%H	12.05%N

EXAMPLE 23

4-Methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent[b]indoi-7-yi methylcarbamate maleate

To a stirred solution of 4-methyl-5-phenylmethylmino-1,2,3.4-tetrahydrocyclopent(b)indol-7-yl methylcarbamate (1.8 g) in acetic acid (25 m) was added sodium cyanoborohydride (0.8 g). The reaction was monitored via TLC and after 2 hours CH₂Cl₂ (50 ml) was added and the solution was washed with saturated NaHCO₂ until neutral. The CH₂Cl₂ layer was dried (Na₂SO₄), filtered and concentrated to give an oil which was purified via flash chromatography, eluting with 21 hexane/acetone. The product-containing fractions were collected and concentrated to an oil which was dissolved in either and thereafter an ethereal maleic acid solution was added until the mixture became acidic. The maleate salt of 4-methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopen(t)b]indo 7-yl methylcarbamate (0.8 grams) which precipitated as a colorless solid was collected.

ANALYSIS:			
Calculated for C ₂₁ H ₂₁ N ₃ O ₂ ·C ₄ H ₄ O ₄	64.51%C	5.85%H	9.03%N
Found	64.13%	5.75%H	8.97%N

EXAMPLE 24

4-Methyl-3-[(2-phenylcyclopropyl)imino]-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate

To a stirred suspension of 1.4-dihydro-7-hydroxy-4-methytcyclopent(b)indol-3(2H)-one (5.0 g) in accionitrile (100 m) was added phenytycycloprotylemine hydrochiodie (4.2 g) followed by triethylamine (2.5 g). The solution was tested at room temperature under a nitrogen atmosphere while titanium (IV) isopropoxide was added in a dropwise manner. The mixture was stirred for 3 hours and thereafter quenched with icalvater. The mixture was filtered, the solids were washed with CH₂CE₂, the layers were separated and the organic portion was dried (Na₂SC₂). After concentration, the crude product was purified via flash chromatography eluting with hexane/acetone (2:1) to give 4-methyl-3-{(2-phenyl-cyclopropy)/impol-1_2.3.4-tetrahydrocyclopropy/impol-1_2.3-tetrahydrocyclopropy/impol-1_2.3.4-tetrahydrocyclopropy/impol-1_2.3.4-tetrahydrocyclopropy/impol-1_2.3.4-tetrahydrocyclopropy/impol-1_2.3.4-tetrahydrocyclopropy/impol-1_2.3.4-tetrahydrocyclopropy/impol-1_2.3-tetrahydrocyclopropy/impol-1_2.3-tetrahydrocyclopropy/impol-1_2.3-tetrahydrocyclopropy/impol-1_2.3-tetrahydrocyclopropy/impol-1_2.3-tetrahydrocyclopropy/impol-1_2.3-tetrahydrocyclopropy/impol-1_2.3-tetrahydrocyclopropy/impol-1_2.3-tetrahy

To a stirred solution of this product (1 0 g) in CH₂C₂ (9 0 ml) was added 1.8-diazabicyclo[5.4.0]undec-7-ene (58 mg) followed by the dropwise addition of methyl isocyanate (0.18 g) in CH₂Cl₂ (1.0 ml). The reaction was monitored via TLC and after 0.5 hour the solution was concentrated and the precipitate was collected and recrystallized twee from acetonitrile to give 4-methyl-3-([2-phenylcyclopropyl)minol-1.2,3.4-tetrahydrocyclopent[b]indol-7-yl methylcar-bamate (0.56 gram).

ANALYSIS:

Calculated for C21H21N3O2: 73.97%C 6.21%H 11.25%N

Found:

73.57%C 6.25%H 11.13%N

EXAMPLE 25

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3-Cyclopropylimino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol

7-Chloroacelyloay-1.4-dillydro-4-methylcyclopent[b]indol-3(2H)-one (15.0 g) and cyclopropylamine (9.6 g) wadre dissolved in 300 ml toluene and cooled to -10°C. Titanium tetrachloride (6.3 g) dissolved in 50 ml toluene was added slowly to the first solution. The reaction mixture was allowed to come up to room temperature and stirred overnight. The next day another 1.5 equivalents of the amine (4.6 g) was added to the reaction mixture and the mixture was stirred for one hour. The reaction mixture was filtered through a pad of slitica gel, eluting with 3:1 hexane/ethyl acetate, giving a yellow oil after removal of solvents. 3-Cyclopropylimino-4-methyl-1,2,3.4-tetrahydrocyclopent[b]-indol-7-oil was isolated as a light yellow solid (3.3 g) after flash chromatography and recrystalization from ethyl acetate.

ANALYSIS:			
Calculated for C ₁₅ H ₁₆ N ₂ O	74.97%C	6.71%H	11.66%N
Found	74.57%C	6.54%H	11.37%N

EXAMPLE 26

3-(N-Cyclopropyl)amino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol maleate

3-Cyclopropylimino-4-methyl-1.2.3.4-tetrahydrocyclopent(b) T-ol (17.3 g) was dissolved in \$1.1 isoprogano/Imstin-anol (250 ml), under N₂ and stirred at room temperature. Sodium borohydride (8.2 g) was added and the reaction mixture was stirred overnight. Thin layer analysis indicated a complete reaction. The solution was cooled to 0°C and water (100 ml) was added slowly. Ethyl acetate (250 ml) was added and, after separating the layers, the organic portion was washed successively with brine (2×1 to 0 ml), and water (2×1 to 0 ml) and dired over Na₂SQ4, and thereafter the solvent was removed <u>in yacuo</u>. The crude material was purified by preparative HPLC using a 2°1 hexane/ethyl acetate solvent system. The free base was isolated as light brownyleption (17.8 g), A stirred solution of the free base (05) in either (200 ml) was treated slowly with a solution prepared from 0.3 g maleic acid, 50 ml Et₂O and 5 m EtO4.

 $3-(N-Cyclopropyl)amino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol maleate was isolated as a light yellow solid (0.8 g) after filtering and drying under <math>N_2$.

ANALYSIS:			
Calculated for C ₁₅ H ₁₈ N ₂ O·C ₄ H ₄ O ₄	63.68%C	6.19%H	7.82%N
Found	63.47%C	6.31%H	7.69%N

EXAMPLE 27

3-(N-cyclopropyl)amino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl phenylmethylcarbonate

3-Cyclopropylamino-4-methyl-1,2.3-4-latrahydrocyclopent[b] indol-7-ol (5.5.9) was dissolved in 250 ml CH₂Cl₂ along with friethylamine (2.8.9) and cooled to 0°C while stirring. Benzyl chloroformate (3.9.9) dissolved in 50 ml CH₂Cl₂ was added slowly to the linst solution. After complete addition, the reaction mixture was allowed to come to room temperature, washed with H₂O (2x 150 ml) and dried over Na₂SCl₂, and the solvent was removed <u>in vacuo</u>. The crude material was unflied by flash column chromatoraph usine ELOAc as the solvent.

3-(N-Cyclopropyl)amino-4-methyl-1,2,3,4-tetrahydro-cyclopent[b]-indol-7-yl phenylmethylcarbonate was isolated as a yellow brown foam (4.1 g).

ANALYSIS:

Calculated for C23H24N2O3: 73.38%C 6.43%H 7.44%N Found: 73.41%C 6.80%H 7.48%N

EXAMPLE 28

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3-(N-Cyclopropyi)amino-1,2,3,3a,4,8b-hexahydro-4-methylcyclopent[b]indol-7-ol

3-Cyclopropylimino-4-methyl-1,2,34-telrahydrocyclopent[b]ndol-7-ol (6.7 g) was placed in a 3-neck flask and cooled to 0°C in an ice-water bath. A 1M solution of borate in 17HF (540 m) was added in a dropwise manner. The mxture was stirred for 1 hour while it was slowly warmed to room temperature. The mixture was cooled back down to 0°C and riffluoreacetic acid (119 mi) was added in a dropwise manner. The solution was stirred for 15 minutes and HFH was removed <u>in vaccor</u> The mixture was neutritazled with 10% NaCH solution, extracted with methylene chloride (4x 500 ml), dried (Na₂SC₄) and concentrated to give 3-(N-cyclopropyl)amino-1,2,3,3a,4,8b-hexahydro-4-methylcy-clopentibilitoble-7-ol (8.8 g).

EXAMPLE 29

3-(N-Cyclopropyl-N-methylaminocarbonyl)amino-1,2,3,3a,4,8b-hexahydro-4-methylcyclopent[b]indol-7-ol hydrochloride

3-(N-Cyclopropyl)amino-1,2,3,3a,4,8b-hexahydro-4-methylcyclopent(blindol-7-ol (8.8 g) was dissolved in CH₂Cl₂ (400 ml) along with triethylamine (4.4 g). The solution was cooled to 0°C and stirred under N₂. Benzyl chloroformate (6.1 g) dissolved in CH₂Cl₂ (50 ml) was added slowly to the first solution. The reaction was monitored by thin layer chromatography while adding an additional equivalent (6.1 g) of the chloroformate until the reaction was complete. The solution was warmed to room temperature before washing with water (2x 100 ml), drying over Na₂SO₄ and concentrating to an oil, which was purified by preparative HPLC using 3:1 hexane/acetone as the solvent system. 3-(N-Cyclopropyl)amino-1,2,3,3a,4,8b-hexahydro-4-methylcyclopent[b]indol-7-yl phenylmethylcarbonate was isolated (7.0 g), which was characterized by NMR, MS and IR. This material was dissolved in CH2Cl2 (250 ml) and the solution treated with 1,8-diazabicyclo[5.4.0]undec-7-ene (DBU; 0.4 g). The mixture was cooled to 0°C and stirred while a solution of methyl isocyanate (1,1 g) in 50 ml CH₂Cl₂ was added slowly. The reaction was monitored by TLC (1:1 hexane/ acetone) while adding another 2.5 equivalents (2.7 g) of methyl isocyanate until the reaction was complete. The solution was warmed to room temperature, washed successively with brine (2x 100 ml) and water (1x 100 ml), dried over Na₂SO₄ and concentrated. The oil was purified by flash column chromatography using ethyl acetate as the solvent 3-(N-Cyclopropyl-N-methylaminocarbonyl)amino-1,2,3,3a,4,8b-hexahydro-4-methylcyclopent[b]indol-7-yl phenylmethylcarbonate was isolated (4.5 g). The material was dissolved in absolute ethanol (190 ml), and 10% palladium on carbon (10% by weight; 0.4 g) was added. The solution was placed in a Parr shaker bottle, charged with Ho (45 psi) and shaken for 2 hours. The catalyst was filtered and the filtrate was concentrated. The oil was triturated with EtOAc (50 ml) and CH₂Cl₂ (5 ml) to give an off-white solid (1.05 g).

3-(N-Cyclopropyl-N-methylaminocarbonyl)amino-1, 2, 3, 34, 4-Bb-hexahydro-4-methylcyclopent(b)indol-7-ol was characterized by NMR. MS and Fi. The solid (0.8 g) was dissolved in 8 1-E₂O/EICH (200 m) initially and eithereal hydrogen chloride was added slowly until the solution became neutral and then more El₂O (800 mi) was added. 3-(N-Cyclopropyl-N-methylamino-carbonyl)amino-1, 2, 3, 34, 4, Bb-hexa-hydro-4-methylcyclopent(b)indol-7-ol hydrochloride was isolated as an off-white solid after filtering and dying under N₂ (0.8 5).

ANALYSIS:			
Calculated for C ₁₇ H ₂₃ N ₂ O ₃ ·HCI	60.44%C	7.16%H	12.44%N
Found	60.77%C	7.41%H	12.67%N

3-(N-Cyclopropyl)amino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate

3-(N-Cyclopropy)Jamino-4-methyl-1.2,3.4-letrahydrocyclopent(blindol-7-ol (2.2.g) was dissolved in CH₂Cl₂ (200 m) with 1,8-diazabicyclo(5.4.0) undec-7-ene (DBU; 0.2.1 g) and the solution was coded to 0°C. A solution of methyl iscoyanate (0.5.2 g in 30 ml CH₂Cl₂) was added slowly to the cooled solution and the reaction was monitored by thin layer chromatography (silica gel. 1:1 hexane/ethyl acetate). After warming to room temperature, the mixture was washed successively with water (2x 100 ml), brine (1x 100 ml) and again with water (1x 100 ml). The organic layer was dried over Na₂SO₄ and the solvent removed in vacuo. The crude material was recrystallized from acetonitrile 3-(N-Cyclopropy)]amino-4-methyl-1.2.3,4-tetrahydrocyclopent(b|indol-7-yl/methylcarbamate was isolated as light yellow/white plates (1.0 o.).

ANALYSIS:			
Calculated for C ₁₇ H ₂₁ N ₃ O ₂	68.21%C	7.07%H	14.04%N
Found	68.08%C	6.57%H	13.97%N

EXAMPLE 31

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1,2,3,3a,4,8b-Hexahydro-4-methyl-3-(N-phenylmethyl-N-methylaminocarbonyl)aminocyclopent[b]indol-7-ylphenylmethylcarbonate

To a stirred solution of 1,2,3,3,4,8,b-hexahydro-4-methyl-5-phenylmethylaminocyclopent(blindol-7-d (12.0 g) in CH₂Cl₂ (50 ml) was added briethylamino (4.0 8 g). The mixture was cooled to 6°C and solution of benzyl chloroformate (8.8 g) in CH₂Cl₂ (50 ml) was added slowly in a dropwise manner. After three hours the reaction mixture was washed with water (2x 100 ml), dried over Na₂So₂ and concentrated to give 17.0 grams of an oil. The crude 4-methyl-3-phenylmethylamino-1,2,3,34,8,4b-axehyldycocyclopen(bliplindol-7-yl) phenylmethylamichorate (17.0 g) was dissolved in CH₂Cl₂ (125 ml) and 1,8-diazabicyclof 6.4 0]undeo-7-ene (0.9 g) was added followed by the dropwise addition of a solution of methyl isocyanate (2.6 g) in CH₂Cl₂ (25 ml). The reaction mixture was stirred for 2 hours and an additional 0.5 gram of methyl isocyanate was added. The reaction mixture was stirred for an additional 15 minutes and threatter concentrated in vacuo to give an oil which was purified by flash chromatography on silica gel elutring with 2:1 hexane/ethyl acetate. The product-containing fractions were collected and concentrated to give an oil (5.5 q).

ANALYSIS:			
Calculated for C22H25N3O4	71.73%C	6.43%H	8.65%N
Found	71.67%C	6.59%H	8.67%N

EXAMPLE 32

1,2,3,3a,4,8b-Hexahydro-4-methyl-3-phenylmethyloxycarbonyl-aminocyclopent[b]indol-7-ol

A solution of 4-methyl-3-phenylmethyllmino-12,34-tetrahydrocyclopentllplindol-7-01 (14.0 g) was placed in a 3-neck flask and cooled to 0°C in an ice-water bath. A solution of 1 M borane/THF in THF (145 m) was added in a dropwise manner. The mixture was stirred for 1 hour while it was slowly warmed to room temperature. The mixture was cooled back to 0°C and trifluoroacetic acid was added in a dropwise manner. The solution was stirred for 15 minutes, neutralized with 10°S NaOH (Ag), extracted with CH₂Cl₂ dried (Na₂SO₄) and concentrated to give 1,2.3,3a, 45-heavilword-4-methyl-3-phenylmethylaminocyclopentiblindio-7-of (14 crams).

20% Palladium hydroxide on carbon (1.4 g) was added to a solution of 1,2,3,3a,4,8b-hexahydro-4-methyl-3-phenylmethylaminocyclopenti[b]indol-7-ol (14 grams) in ethanol (100 ml) and the mixture was hydrogenated at 45 psi H₂ pressure using a Parr apparatus at 50°C for 5 hours. The mixture was filtered and the solution was concentrated to give 3-amino-1,2,3,3a,4,8b-hexahydro-4-methyloxiclopentibilindol-7-ol (10,7 grams).

To a solution of 3-amino-1,2,3.3.4, 8b-haxahydro-4-methylcyclopent[b]indol-7-ol (10.7 grams) in methylene chloride (12.5 ml) was added triethylamine (5.6 grams) followed by the dropwise addition of benzyl chloroformate (10.0 grams) in methylene chloride (25 ml). The mixture was stirred for 2 hours, extracted with water, dried (Na₂SQ₄) and concentrated The product was purified by chromatography on silica gel, eluting with 2.1 hexane/acetone to provide 1,2.3 a.4 8b-kavahydro-4-methyl-3-phenylmethyloxycarbonylaminocyclopent[blindol-7-ol.

1,2,3,3a,4,8b-Hexahydro-4-methyl-3-(N-phenylmethyloxycarbonyl)aminocyclopent[b]indol-7-ylmethylcarbamate

To a stirred solution of 1,2,3,3.4 (Bb-hexatlydro-4-methyl-3-(N-phenylmethyloxycarbonyl)aminocyclopent[b]indoi-7-ol (1.8 g) in CH₂Cl₂ (75 m) was added 1,8-diazabicyclo[5,4.0]undec-7-ene (0.12 g) followed by the dropwise addition of a solution of methyl isocycarate (0.36 g) in CH₂Cl₂ (25 m)). The reaction mixture was stirred for 2 hours and an additional 0.1 gram of methyl isocycarate was added. The reaction mixture was stirred for an additional 15 minutes and concentrated in *vacuo* to give an oil which was purified by flash-chromategraphy on silicia gel eluting with 2 1 hexane/ ethyl acetate. The product which crystallized from the pure fractions was collected by filtration to give 600 mg and the filtrate was concentrated to give an oil (800 mg) which crystallized upon standing.

ANALYSIS:			
Calculated for C22H25N3O4	66.82%C	6.37%H	10.63%N
Found	66.91%C	6.47%H	10.66%N

EXAMPLE 34

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3-Methylaminocarbonyloximino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate hemihydrate

To a stirred suspension of 3-hydroxyimino-4-methyl-12,3.4-tetrahydrocyclopent[b]indol-7-ol (3.0 g) in CH₂Cl₂ (100 m) was added 1,8-diazabicyclofs, 4.0 pluder-7-ene (630 mg) followed by methyl iscoyanate (1.9 g) and the miture was stirred overnight at ambient temperature. The mixture was concentrated in vacuo and the resulting solid was recrystallized from ethanol to give 3-methylaminocarbonyloximino-4-methyl-1,2,3.4-tetrahydrocyclopent[b]indol-7-yl methylcarbonate hemitydrate (1.7 g).

ANALYSIS:			
Calculated for C ₁₆ H ₁₈ N ₄ O ₄ -0.5H ₂ O	56.68%C	5.66%H	16.53%N
Found	56.57%C	5.46%H	16.68%N

35 EXAMPLE 35

1,2,3,3a,4,8b-Hexahydro-4-methyl-3-methylaminocarbonyl-aminocyclopent[b]Indol-7-ol hydrochloride monohydrate

A solution of 1,2,3,3,4,8b-hexahydro-4-methyl-3-(N-phenymethyl-N-methylaminocarbonyl)aminocyclopont[bjin-dol-7-yi phenylmethyl carbonate (1,7,9) in glacial acetia cald (1,0 m) was hydrogenated at 45 ps i h_s and 50°C in the presence of 20°S Fd hydroxide on carbon utilizing a Parr apparatus. After four hours, TLC indicated a compilete reaction with the formation of a major product as well as a side product. The Pd catalyst was filtered under nitrogen and the filtrate concentrated in vazuo. The material was chromatographed on silica gel situling with 10°S. McCH/CH₂Cb₂. The product-containing fractions were collected and concentrated. The resulting oil was dissolved in EiOH (25 ml) and Ego (150 ml), the solution was filtered, and ethereal HCI was added to the filtrate until the solution became acidic. The coloriass solid which formed was collected under Ng and dried under vacuum to give 1,2,3,3a,4,8b hexahydro-4-methyl-3-methylaminocatopolyamino cyclopenel[bjindot-7-ol hydrochloride monohydrate (0.25 ml).

ANALYSIS:			
Calculated for C ₁₄ H ₁₉ N ₃ O ₂ -HCl-H ₂ O	53.25%C	7.02%H	13.31%N
Found	53.26%C	6.54%H	12.78%N

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1,2,3,3a,4,8b-Hexahydro-4-methyl-3-(N-phenylmethyl-N-methylaminocarbonyl)aminocyclopent[b]indol-7-ol hydrochloride

A solution of 1,2,3,3,4,8b-hexahydro-4-methyl-3-(N-phenylmethyl-N-methylaminocarbonyl)aminocyclopent[b]in-dol-7-yl phenylmethyl carbonate (2.0 g) in absolute ethanol (100 ml) was hydrogenated at 45 psi l4, n the presence of 5% Pd-acrbon utilizing a Parr apparatus. After two hours, TLC indicated a complete reaction with the formation of a single product. The Pd catalyst was filtered under nitrogen and the filtrate concentrated in vacuo. The resulting of was dissolved in EIOAe (25 ml) and El₂O (150 ml), the solution was filtered and ethereal HCI was added to the filter until the solution became acidic. The colorless solid which formed was collected under N₂ and dried overnight at 40°C under vacuum to give 1,2,3,3a,4,8b-haxshydro-4-methyl-3-(N-phenylmethyl-N-methylaminocarbonyl)aminocyclopent [blindof-7-ol hydrochloride (1,4 o).

ANALYSIS:			
Calculated for C21H25N3O2+HCI	65.02%C	6.76%H	10.83%N
Found	64.95%C	6.85%H	10.84%N

20 EXAMPLE 37

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4-Methyl-3-phenylethyliming-1,2,3,4-tetrahydrocyclopent[b]Indol-7-ol hemihydrate

To a stirred solution of 7-hydroxy.4-methyl-1,4-dihydrocyclopent(b)-3-one (5.0 g) in acetonitrile (100 m)) were added phenethylamine (6.0 g) and titanium isopropoxide (14.1 g), and the resulting mixture was stirred under nitrogen at ambient temperature for 3 hours. The mixture was poured onto ice/water (200 mt) and thereafter, CH₂Cl₂ (500 mt) was added. The mixture was filtered, and the organic layer was separated from the filtrate, dried over sodium sulfate and concentrated in vacuo. Crystallization from CH₂Cl₂hexane provided 4-methyl-3-phenylethylimino-1,2,3,4-tetrahydro-cycloomtibilidol-7-ot heminytrate (3.0 q).

ANALYSIS:			
Calculated for :C ₂₀ H ₂₀ N ₂ O 1/2H ₂ O	76.65%C	6.75%H	8.94%N
Found	76.53%C	6.38%H	8.89%N

EXAMPLE 38

4-Methyl-3-phenylethylimino-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate

To a stirred solution of 4-methyl-3-phenylethylimino-1,2,3,4-tetrahydrocyclopent[b]indoi-7-ol hemihydrate (1.43 g) in CH₂Cl₂ (25 mi) was added 1,8-diazabicyclof5,4.0]undeo-7-ene (0.11 g). Methylisocyanate (0.27) in CH₂Cl₂ (20 mi) was added to the reaction mixture. The reaction was monitored by TLC and after 3 hours the CH₂Cl₂ was evaporated off. The brown residue was recrystallized from acetonlinie.

ANALYSIS:			
Calculated for : C ₂₂ H ₂₃ N ₃ O ₂	73.11%C	6.41%H	11.63%N
Found			11.65%N

50 EXAMPLE 39

4-Methyl-3-(2-phenylethyl)amino-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate hydrochloride hemihydrate

To a stirred solution of 4-methyl-3-(2-phenylethyl)imino-1,2,3,4-tetrahydrocyclopent[b]indol-7-methyl carbamate (0.80 g) in acetic acid (8 mt), shanol (8 mt), isopropanol (8 mt) and tetrahydrofuran (8 mt) was added sodium cyanoborohydride (0.35 g) under nitrogen. The reaction was monitored by TLC and after 1 hour the solution was neutralized with saturated NaHOOs, extracted with EtOAc, dried over NasSOs, and concentrated in vacuo. The resulting

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yellow oil was dissolved in a minimum amount of EIOAc, diluted with ether and thereafter, an etheral HCl solution was added. The resulting white solid was collected via filtration. Crystallization from ethanol afforded 4-methyl-3-(2-phenylethyl)amino-1,2,3,4-tetrahydrocyclopen([b]indo-17-yl methyl carbamate hydrochloride hemihydrate (0.68 g). The reaction was repeated and the material combined.

ANALYSIS:			
Calculated for C ₂₂ H ₂₅ N ₃ O ₂ -HCI-½H ₂ O	64.62%C	6.65%H	10.28%N
Found	64.56%C	6.72%H	9.91%N

EXAMPLE 40

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4-Methyl-3-(2-propynyl)imino 1,2,3,4-tetrahydrocyclopent[b]indol-7-yl methylcarbamate

To a stirred suspension of 7-hydroxy-4-methyl-1,4-dihydrocyclopent[b]indol-3-(2H)-one (5.5 g) in acetonitrile (100 m) was added propargyl amine (3.0 g), the solution was stirred at room temperature under a ritrogen atmosphere while titanium (iv) sporpoxible (15.6 g) was added in a dropwise manner. The mixture was stirred for 16 hours before quenching with ice water. The mixture was filtered, the solids were washed with CH₂Cl₂, the layers were separated and the organic portion was dried (Nag-SQ₄). After concentration, the crude product was purified via flash chromatography eluting with hexane/scatone (2:1) to give 4-methyl-3-(2-propynyllimino-1,3-4-tetrahydrocyclopent bi indol-7.

To a stirred solution of 4-methyl-9-(2-propynyl)rinno-1,2,3.4-tetrahydrocyclopent(b)indol-7-ol (3.4 g) in CH₂Cl₂ (15.0 ml) was added 1.8-diazabicyclo[5.4.0]undec-7-ene (326 mg) followed by the dropwise addition of methyl isocyanate (0.8 g) in CH₂Cl₂ (5.0 ml). The reaction was monitored via TLC and after 1.0 hour, the solution was concentrated and the crude product was purified via flash chromatography elluting with hexane/acetone (2:1). The product which precipitated out of the pure fractions was collected to give 4-methyl-3-(2-propynyl)imino-1,2,3,4-tetrahydrocyclopent (bilindol-7-vi methydrachamate 1.2 grams) and the fractions were concentrated to give a methydroadbard of the grams and the fractions were concentrated to give an additional 0.9 gram.

ANALYSIS:

50	Calculated for C ₁₇ H ₁₇ N ₃ O ₂ :	69.14%C	5.80%H	14.23%N
35	Found:	68.94 % C	5.81%H	13.94%N

EXAMPLE 41

4-Methyl-3-(2-propynyi)amino-1,2,3,4-tetrahydrocyclopent[b]indoi-7-yi methylcarbamate hydrochloride monohydrate

To a stirred solution of 4-methyl-S-(2-propynyllymino-1,2.3.4-tetrahydrocyclopent[b]inded-T-yf methylacabarnate (1: 1g) in acetic acid (10 ml) was added sodium cyanoborohydride (0.57 g). The reaction was monitored via TLC and after 2 hours, methylene chiloride (50 ml) was added and the solution was washed with saturated NaHCO₂ until neutral. The methylene chiloride layer was dried (Na₂SO₄), filtered and concentrated. The resulting material was chromatographed on sillea get, eluting with 2: 1 hexane/acetore and the pure fractions were collected and concentrated. The resulting solid was dissolved in a minimum amount of E(OAc, diluted with ether and thereafter, ethereal HCi solution was added The resulting solid was collected via filtration under nitrogen to give 4-methyl-8-2-propynyllacin-1,2.3.4-tetrahydrocyclopent[b]indol-7-yf methylcarbarnate hydrochloride monchydrate (0.4 grams). The reaction was repeated and the products were combined.

ANALYSIS:

Calculated for C₁₇H₁₉N₃O₂·HCl·H₂O: 58.04%C 6.30%H 11.94%N

Found: 57.99%C 6.03%H 11 R2%N

EXAMPLE 42

4-Methyl-3-(2-phenylethyl)amino-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl benzylcarbamate hydrochloride hemihydrate

To a stirred solution of 4-methyl-3-(2-phenylethyl)imino-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol (2 70 g) in CH₂Cl₂ (100 ml) was added 1,8-diazabicyclo[5,4.0]undecene (0.21 g). Benzyl isocyanate (0.83 g) was added to the reaction mixture via a syringe and the mixture stirred under nitrogen. Additional benzyl isocyanate was added after 120 and 180 minutes in 1/4 and 1/2 equivalents, respectively. The reaction was monitored by TLC and after 185 minutes the solution was concentrated in vacuo. The crude reaction residue (2.72 g) showed carbamate formation according to proton NMR and MS. The residue was dissolved in glacial acetic acid (75 ml) with stirring under nitrogen. A yellow precipitate formed upon addition of sodium cyanoborohydride (0.98g) and dissolved after 30 minutes. One equivalent of sodium cvanoborohydride was added after 3 hours. After 30 minutes, TLC showed complete reaction. The reaction mixture was neutralized with saturated sodium bicarbonate solution, extracted with ethyl acetate, dried over Na₂SO₄, and concentrated in vacuo. The free base was dissolved in ether and an ethereal solution of HCI was added. The resulting white solid was collected via filtration.

ANALYSIS:			
Calculated for C ₂₈ H ₃₀ N ₃ O ₂ ·HCI·½H ₂ O	69.34%C	6.44%H	8.66%N
Found	69.45%C	6.30%H	8.72%N

EXAMPLE 43

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4-Methyl-3-[2-(4-morpholinyl)ethyl]imino-1,2,3,4-tetrahydrocyclopent[b]indol-7-ol

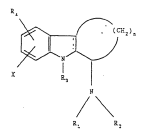
To a stirred solution of 7-hydroxy-4-methyl-1,4-dihydrocyclopent[b]indol-3-one (8.00 q) in acetonitrile (125 ml) under nitrogen were added 4-(2-aminoethyl)morpholine (10.35 g) and titanium isopropoxide (22.60 g). The reaction was monitored by TLC and after two hours additional equivalents of 4-(2-aminoethyl)morpholine (5.17 g) and titanium isopropoxide (11.30 g) were added. Fourteen hours later the reaction was quenched with water (200 ml). EtOAc (200 ml) was added and the mixture stirred for fifteen minutes and filtered. The layers were separated, the aqueous layer was extracted with ethyl acetate, and the combined organics were dried (Na₂SO₄) and concentrated in vacuo. The resulting vellow solid was dried vielding 6.65 a. of product. A 2 a sample of the solid was further purified by crystallization from CH₂Cl₂/hexane to afford 1.2 g of 4-methyl-3-(2-morpholinoethyl)imino-1.2.3.4-tetrahydrocyclopent[b]indol-7-ol.

ANALYSIS:			
Calculated for C ₁₈ H ₂₃ N ₃ O ₂	68.98%C	7.40%H	13.41%N
Found	68.78%C	7.52%H	13.26%N

45 Claims

Claims for the following Contracting States: AT, BE, CH, DE, DK, FR, GB, IT, LI, LU, NL, PT, SE

1. A compound of the formula,



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n is 2, 3, 4 or 5;

X is hydrogen, C1-C6-alkyl, C1-C6-alkoxy, hydroxy, halogen, trifluoromethyl or nitro;

 R_1 is hydrogen, C_1 - C_6 -alkyl, C_2 - C_6 -alkyn, C_3 - C_6 -alkyn, R_1 -alkyl, R_1 - R_2 -alkyl, R_1 - R_2 -alkyl, R_2 - R_3 -alkyl, R_3 - R_4 - R_4 - R_4 - R_5 - R_5 - R_5 - R_6 -alkyl, R_3 - R_5 - R_6 -R

the group "Alk" signifying a divalent C_1 - C_6 -alkylene group, and Y signifying hydrogen, C_1 - C_6 -alkyl, phenyl or phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above, C_1 - C_6 -alkyl, formyl, C_1 - C_6 -alkyl, formyl, C_1 - C_6 -alkylaminocarbonyl, benyl-polycoyarbonyl or C_1 - C_6 -alkylaminocarbonyl; or

n₂ is rivologen, 01-06-aikyr, formyr, 01-06-aikyrcarbonyr, benzyloxycarbonyr or 01-06-aikyraminocarbonyr, or alternatively, the group



as a whole is

$$-N$$
 $-N$ $-N$

$$-N$$
 NH $-N$ N- C_1 - C_6 -alkyl

wherein the phenyl group may be substituted as indicated above,

 H_3 is hydrogen, C_1 - C_6 -alkyl, phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above, C_1 - C_6 -alkylcarbonyl or C_1 - C_6 -alkoxycarbonyl; H_2 is hydrogen, CH_3 .

Oſ

wherein

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 $H_5 \text{ is } C_1 - C_6 \text{-}alkyl, C_2 - C_6 \text{-}alkenyl, C_3 - C_6 \text{-}alkynyl, C_3 - C_7 \text{-}cycloalkyl, C_3 - C_7 \text{-}cycloalkyl, C_6 - C_6 \text{-}alkyl, phenyl, phenyl, C_3 - C_7 \text{-}cycloalkyl, wherein the phenyl group may be substituted as indicated above; and the phenyl group may be substituted as ind$

R₆ is hydrogen, C₁-C₆-alkyl, phenyl or phenyl-C₁-C₆-alkyl, wherein the phenyl group may be substituted as indicated above;

or alternatively the group

$$-N$$
 R_{ϵ}

as a whole is

$$N - N$$
 $N - N$ $N - C_1 - C_6$ alkyl

$$N-\text{phenyl or}$$
 $N-\text{phenyl or}$ $N-\text{phenyl or}$ $N-\text{C}_1\text{-C}_6$ alky

wherein the phenyl group may be substituted as indicated above,

and

 R_7 is C_1 - C_6 -alkyl, phenyl or phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above:

with the proviso that R₄ is not hydrogen or hydroxy, when n is 4 or 5; or a pharmaceutically acceptable acid addition salt thereof.

- A compound as defined in claim 1, where n is 3.
- 10 3. A compound as defined in claim 2, where
 - X is hydrogen or hydroxy.

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- P1 is hydrogen, C3-C6-alkynyl, C3-C7-cycloalkyl, phenyl or phenyl-C1-C6-alkyl,
- R₂ is hydrogen, formyl, benzyloxycarbonyl or C₁-C₆-alkylaminocarbonyl,
- R₃ is hydrogen or C₁-C₆-alkyl,
- R4 is hydrogen or a group of the formulae

wherein R_5 is C_1 - C_6 -alkyl or phenyl- C_1 - C_6 -alkyl and R_6 is hydrogen, and R_7 is phenyl- C_1 - C_6 -alkyl, where each phenyl group in the definitions of R_1 , R_5 and R_6 may be substituted as indicated in claim 1.

- 4. A compound as defined in claim 3, where
 - X is hydrogen
 - R₁ is C₃-C₇-cycloalkyl, C₃-C₆-alkynyl, phenyl-C₃-C₇-cycloalkyl or phenyl-C₁-C₆-alkyl, wherein the phenyl group may be substituted as indicated in claim 1
 - Ro is hydrogen
 - R₄ is hydrogen or a group of the formula



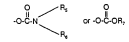
where R₅ is C₁-C₆-alkyl and R₆ is hydrogen.

- The compound as defined in claim 1, which is 3-cyclopropylamino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]-indol-7-yl methylcarbonate.
- The compound as defined in claim 1, which is 4-methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent[b]-indol-7-yl methylcarbonate.
 - The compound as defined in claim 1, which is 1,2,3,4-tetrahydro-cyclopent[b]indol-3-(2-propynyl)amine.
- A compound of the formula III
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EP 0 496 314 B1

where H₃, F₄, X and n are as defined in claim 1, and H₈ is hydroxy, C₁-C₆-alkoxy, amino-C₁-C₆-alkoxy, C₁-C₆-alkoxy, C₂-C₇-cycloalky, C₃-C₇-cycloalky, herein the phenyl group may be substituted as indicated in claim 1, C₁-C₆-alkylcarbonyloxy or C₁-C₆-alkylaminocarbonyloxy or a pharmaceutically acceptable addition sall thereof.

- 9. A compound as defined in claim 8, where n is 3,
- 10. A compound as defined in claim 9 where
 - X is hydrogen, hydroxy or C₁-C₆-alkoxy
 - R₃ is hydrogen or C₁-C₆-alkyl
- R₄ is hydrogen, a group of the formula



- wherein R₅, is C₁-C₆-alkyl or phenyl-C₁-C₆-alkyl, R₆ is hydrogen, and R₇ is C₁-C₆-alkyl or phenyl-C₁-C₆-alkyl.

 R6 is hydroxy, G₂-G₂-alkynyl, amino-C₁-C₆-alkoxy, C₁-C₆-alkylcarbonyloxy, C₁-C₆-alkylaminocarbonyloxy, C₃-C₇-cycloalkyl, phenyl-C₃-C₇-cycloalkyl, phenyl-C₃-G₇-Cycloalkyl, wherein each phenyl group in the definitions of R₅, R₆ and R₃ may be substituted as indicated in claim 1.
- The compound as defined in claim 8, which is 4-methyl-3-phenylmethylimono-1,2,3,4-tetrahydrocyclopent[b]-indol 7-ol.
 - A pharmaceutical composition which comprises as the active ingredient a compound as defined in claims 1 or 8 and a suitable carrier therefor.
- 45 13. Use of a compound as defined in claim 1 for the preparation of a medicament having memory dysfunction alleviating and/or antidepressant activity.
 - 14. Use of a compound as defined in claim 8 for the preparation of a medicament having antidepressant activity.
- 50 15. A process for the preparation of a compound as defined in claim 1, which comprises
 - a) reducing a compound of the formula XVI

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(XVI)

where R_9 , X and n are as defined in claim 1 and R_{12} is hydrogen, methoxy or hydroxy to form a compound of the formula I, where R_9 , X and n are as defined, R_4 is hydrogen, methoxy or hydroxy, and R_1 and R_2 are hydrogen, or

b) reacting a compound of the formula XV

where R_3 , X and n are as defined in claim 1 and R_{12} is hydrogen, methoxy or hydroxy, with titanium isopropoxide and a compound of the formula

where the group

is

$$NH - N - C_1 - C_6$$
 alkyl

— N N-phenyl or —N
$$N-C_1-C_6$$
 alkylphenyl

wherein the phenyl group may be substituted as indicated in claim 1,

followed by reduction with sodiumborohydride to form a compound of the formula I, wherein $\rm H_3$, X and n are as defined, $\rm H_4$ is as defined for $\rm H_{12}$ above and the group

as a whole has the meaning given for

above, or

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c) reducing a compound of the formula XVIII

(XVIII)

where H_3 , X and n are as defined in claim 1, H_{12} is hydrogen, methoxy or hydroxy, and H_{14} is C_1 - C_6 -alkyl, C_2 -

 C_6 -alkenyl, C_3 - C_6 -alkynyl, C_3 - C_7 -cycloalkyl, C_3 - C_7 -cycloalkyl, phenyl- C_1 - C_6 -alkyl, phenyl- C_3 - C_7 -cycloalkyl or phenyl, wherein the phenyl group may be substituted as indicated in claim 1, to form a compound of the formula I, where R_3 . X and n are as defined in claim 1, R_{12} is as defined above, R_2 is hydrogen and R_1 is as defined for R_1 above.

d) optionally reducing a compound of the formula I, wherein R_0 , R_4 X and n are as defined in claim 1 and R_1 and R_2 are hydrogen with the aid of borane/letrahydrofuran and trifluoroacetic acid to form a compound of the formula ia

$$\begin{array}{c} \mathbb{R}_4 \\ \mathbb{R}_4 \\ \mathbb{R}_3 \end{array} \qquad \begin{array}{c} (\mathsf{CH}_2)_n \\ \mathbb{NH}_2 \end{array}$$

where R₂, R₄, X and n are as defined,

e) optionally reacting a compound of the formula I, where B₃, B₄, X and n are as defined in claim 1 and B₁ and B₃ are hydrogen, with a compound of the formula Hal B₁₅, where B₁₅ is C₁-C₅-alkyl, C₂-C₅-alkenyl, C₃-C₅-alkyl, N₂-C₇-C₅-alkyl, wherein the phenyl group may be substituted as indicated in claim 1, or a group of the formula

or

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where Alk and Y are as defined, to form a group of the formula I, where R₃, R₄, X and n are as defined, R₁ has the meaning of R₁₅ as defined above, and R₂ is hydrogen,

1) optionally reacting a compound of the formula I, where R₀, R₄, X and n are as defined in claim 1, R₁ is hydrogen, C₁-C₂-alkyl, C₂-C₇-cycloalkyl, N₂-C₇-cycloalkyl, N₂-C₇-cycloalkyl, hydrein the phenyl group may be substituted as indicated in claim 1, and T₂ is hydrogen, with formic acid, to form a compound of the formula I, where R₃, R₄, X and n are as defined in claim 1, R₁ is a defined above and R₂ is formyl.

g) optionally reacting a compound of the formula I where R_3 , R_4 , X and n are as defined in claim 1, R_1 , is hydrogen, C_1 - C_6 -allkyl, C_2 - C_6 -allkenyl, C_3 - C_7 -cycloalkyl, C_3 - C_7 -cycloalkyl, therein the phenyl group may be substituted as indicated in claim 1, and R_2 is hydrogen, with an acyl chloride of the formula R_1 - C_6 -Col where R_1 - C_6 -alkyl, to form a compound of the formula 1, where R_3 , R_4 , X and n are as defined in claim 1, R_1 is as defined above and R_2 is C_1 - C_6 -alkyl, and C_1 - C_1

h) optionally reacting a compound of the formula I, where Pa, Ra, X and n are as defined in claim 1 with the

proviso that R_4 is not hydroxy, R_1 is hydrogen, C_1 - C_6 -alkyl, C_2 - C_6 -alkyl, C_3 - C_6 -alkyl, C_3 - C_6 -alkyl, C_3 - C_7 -cycloalkenyl, phenyl- C_1 - C_6 -alkyl or phenyl- C_3 - C_7 -cycloalkyl, wherein the phenyl group may be substituted as indicated in claim 1, and R_2 is hydrogen, with an benzylchloroformate to form a compound of the formula I, where R_3 , R_4 , and n are as defined above, R_4 is as defined above and R_3 is benzyloxy carbonyl.



where R₁₇ is as defined,

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j) optionally reacting a compound of the formula I, where R_1 , R_2 , R_3 , X and n are as defined in claim 1 and R_4 is hydroxy with the proviso that R_2 is not C_1 - C_6 -alkylaminocarbonyl, and with a chloroformate of the formula



where R_7 is as defined in claim 1, to form a compound of the formula I, where R_1 , R_2 , R_3 , X and n are as defined above and R_2 is the group



where R₇ is as defined in claim 1.

k) optionally reacting a compound of the formula I, where R_1 , R_3 , X and n are as defined in claim 1, R_2 is hydrogen and R_4 is the group



where R_7 is benzyl, with an isocyanate of the formula R_{17} -N=C=O to form a compound of the formula I, where R_1 , R_2 , R_3 , R_4 , X and n are as defined above, and R_2 is the group



where R₁₇ is C₁-C₆-alkyl, phenyl or phenyl-C₁-C₆-alkyl, wherein the phenyl group may be substituted as indicated in claim 1.

I) optionally reacting a compound of the formula I, where R_1 , R_2 , R_3 , X and n are as defined in claim 1 and R_4 is hydroxy, with a compound of the formula R_{17} N=C=O where R_{17} is C_1 - C_6 -allkyl, phenyl or phenyl- C_1 - C_6 -allkyl, where in the phenyl group may be substituted as indicated in claim 1, to form a compound of the formula I, where R_1 , R_2 , R_3 , X and R_4 are defined above and R_4 is the group



where R₁₇ is as defined above.

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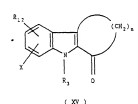
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- 25 16. A process for the preparation of a compound of the formula III as defined in claim 8, which comprises
 - a) reacting a compound of the formula XV



where R_3 X and n are as defined in claim 1, and R_{12} is hydrogen, hydroxy or methoxy, with hydroxylaminhydrochloride to form a compound of the formula III where R_3 . X and n are as defined above, R_4 has the meaning of R_2 above, and R_3 is hydroxy.

b) optionally reacting a compound of the formula III, where R_3 , X and n are as defined in claim 1, R_4 is hydrogen, hydroxy or methoxy and R_3 is hydrogen with a compound of the formula $Br-R_{13}$ *Ni R_2 , where R_{13} is $C_1\cdot C_5$ * alkylene, to form a compound of the formula III where R_3 , R_{12} , X and n are as defined and R_3 is amino- C_1 * C_5 -alkoxy, or

c) reacting a compound of the formula XV

where B_0 X and n are as defined in claim 1 and B_{12} is hydrogen, hydroxy or methoxy, with an amine of the formula NH_2B_{14} where B_{14} is C_1-C_2 -alkyl, C_2-C_2 -alkynyl, C_3-C_2 -cycloalkyl, C_3-C_2 -cycloalkyl, phenyl- C_3-C_2 -cycloalkyl, or phenyl, wherein the phenyl group may be substituted as indicated in claim 1, to form a compound of the formula III, where B_0 X and n are as defined above B_0 has the meaning of B_{12} above, and B_1 has the meaning of B_{12} above, and B_1 has the meaning of B_{12} above, and B_2 has the meaning of B_{12} above, and B_3 has the meaning of B_{12} above.

d) optionally reacting a compound of the formula III, where R_3 , X and n are as defined in claim 1, R_4 is hydrogen, or methroxy and R_3 is hydroxy, with an isocyanate of the formula R_{17} -NeC=O where R_{17} is C_1-C_6 -alkyl, herein the phenyl group may be substituted as indicated in claim 1, to form a compound of the formula III, where R_3 , R_4 , X and n are as defined above and R_8 is the group



where R₁₇ is as defined.

e) optionally reacting a compound of the formula III, where R_8 , X and n are as defined in claim 1, R_4 is hydrogen, hydroxy or methoxy and R_8 is hydroxy, with an acylchloride of the formula



or an acyl anhydride of the formula $(R_1-CO)_2O$ where R_1 , $r_1 = C_1-C_6$ -alkyl, phenyl or phenyl- C_1-C_6 -alkyl, wherein the phenyl group may be substituted as indicated in claim 1, to form a compound of the formula III, where R_1 , R_2 , X and n are as defined above and R_3 is the group.



where R₁₇ is as defined above,

17. A compound of the formula II

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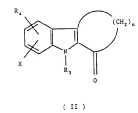
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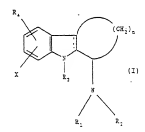
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20 where R₃, R₄, X and n are as defined in claim 1.

Claims for the following Contracting States: ES, GR

25 1. A process for the preparation of a compound of the formula I,



where

n is 2, 3, 4 or 5;

X is hydrogen, C1-C6-alkyl, C1-C6-alkoxy, hydroxy, halogen, trifluoromethyl or nitro;

$$\label{eq:continuous} \begin{split} &R_1 \text{ is hydrogen. } C_1\text{-}C_8\text{-}alkyl, C_2\text{-}C_8\text{-}alkynl, C_3\text{-}C_8\text{-}alkylm, lamino-}C_1\text{-}C_8\text{-}alkyl, C_1\text{-}C_8\text{-}alkylamino-}C_1\text{-}C_8\text{-}alkyl, C_3\text{-}C_7\text{-}cycloalkyl, C_3\text{-}C_7$$

$$-$$
 Alk $-$ N $-$ Alk $-$ N $-$ Alk $-$ N $-$ O

the group "Alk" signifying a divalent C_1 - C_6 -alkylene group, and Y signifying hydrogen, C_1 - C_6 -alkyl, phenyl or phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above; P_2 is hydrogen, C_1 - C_6 -alkyl, formyl, C_1 - C_6 -alkylcarbonyl, benzyloxycarbonyl or C_1 - C_6 -alkylaminocarbonyl; or alternatively, the group



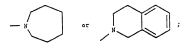
as a whole is

$$-N$$
 NH $-N$ N-C₁-C₆-alkyl

or

$$-N$$
 $N-C_1-C_6$ alky1pheny1

wherein the phenyl group may be substituted as indicated above,



 R_0 is hydrogen, $C_1 - C_6$ -alkyl, phenyl- $C_1 - C_6$ -alkyl, wherein the phenyl group may be substituted as indicated above, $C_1 - C_6$ -alkoxycarbonyl, R_0 is hydrogen, -OH,

or

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wherei

$$\label{eq:problem} \begin{split} &R_{5}\text{ is }C_{1}-C_{6}\text{-alkyl},\ C_{2}-C_{6}\text{-alkenyl},\ C_{3}-C_{6}\text{-alkenyl},\ C_{3}-C_{7}\text{-cycloalkyl},\ C_{3}-C_{7}\text{-cycloalkyl},\ C_{3}-C_{7}\text{-cycloalkyl},\ c_{1}-C_{6}\text{-alkyl},\ c_{2}-C_{6}\text{-alkyl},\ c_{2}-C_{7}\text{-cycloalkyl},\ c_{3}-C_{7}\text{-cycloalkyl},\ c_{3}-C_{7}\text{-c$$

 R_6 is hydrogen, C_1 - C_6 -alkyl, phenyl or phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above; or alternatively the group



as a whole is

$$-N$$
 NH $-N$ N-C₁-C₆-alkyl

wherein the phenyl group may be substituted as indicated above,

and

 R_7 is $C_1\text{-}C_8$ -alkyl, phenyl or phenyl- $C_1\text{-}C_6$ -alkyl, wherein the phenyl group may be substituted as indicated above,

with the proviso that R_4 is not hydrogen or hydroxy, when n is 4 or 5; or a pharmaceutically acceptable acid addition salt thereof, which comprises

a) reducing a compound of the formula XVI

(XVI)

where B_3 , X and n are as defined in claim 1 and B_{12} is hydrogen, methoxy or hydroxy, to form a compound of the formula I, where B_3 , X and n are as defined, B_4 is hydrogen, methoxy or hydroxy, and B_1 and B_2 are hydrogen, or

b) reacting a compound of the formula XV

where $\rm R_3$, X and n are as defined in claim 1 and $\rm R_{12}$ is hydrogen, methoxy or hydroxy, with titanium isopropoxide and a compound of the formula

where the group

$$\binom{z}{z}$$

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45 -N N-phenyl or -N N-
$$C_1$$
- C_6 -alkyl N- C_1 - C_6 - C_6 -alkyl N- C_1 - C_6

wherein the phenyl group may be substituted as indicated above,



followed by reduction with sodiumborohydrode to form a compound of the formula I, wherein R₃, X and n are as defined, R₄ is as defined for R₁₂ above and the group

as a whole has the meaning given for

above, or

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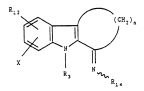
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c) reducing a compound of the formula XVIII



(XVIII)

where R_3 , X and n are as defined in claim 1, R_{12} is hydrogen, methoxy or hydroxy, and R_{14} is $C_1 \cdot C_2 \cdot e$ alkonyl, $C_3 \cdot C_2 \cdot c$ coloalkyn, $C_3 \cdot C_3 \cdot c$ coloalkyn, or phenyl, wherein the phenyl group may be substituted as indicated above, to form a compound of the formula $I_3 \cdot I_3 \cdot c$ color $I_3 \cdot$

d) optionally reducing a compound of the formula I, wherein R_3 , R_4 , X and n are as defined in claim 1 and R_1 and R_2 are hydrogen with the aid of borane/letrahydrofuran and trifluoroacetic acid to form a compound of the formula ia

where R₃, R₄, X and n are as defined,

e) optionally reacting a compound of the formula I, where R_3 , R_4 , X and n are as defined in claim 1 and R_1 and R_2 are hydrogen, with a compound of the formula Hall R_{15} , where R_{15} is C_7 - C_6 -alkyl, C_2 - C_6 -alkenyl, C_3 - C_6 -alkyl, phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above, or a group of the formula

or

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where Alk and Y are as defined, to form a group of the formula I, where R₃, R₄, X and n are as defined, R₁ has the meaning of R₁₅ as defined above, and R₂ is hydrogen,

I) optionally reacting a compound of the formula I, where R_0 , R_0 , X and n are as defined in claim 1, R_1 is hydrogen, C_1 - C_2 -callkyl, C_2 - C_3 -callkynyl, C_3 - C_3 -cycloalkyl, C_3 - C_3 -cycloalkyl, heneyl- C_3 - C_3 -callkyl or phenyl- C_3 - C_3 -cycloalkyl, wherein the phenyl group may be substituted as indicated above, and R_2 is hydrogen, with formic acid, to form a compound of the formula I, where R_3 , R_4 , X and n are as defined in claim 1, R_1 , is as defined above and R_3 is C_3 - C_3 - C_4 - C_4 - C_5 - $C_$

g) optionally reacting a compound of the formula I where R_b , R_a , X and n are as defined R_i is hydrogen, C_1 - C_6 -alkyl, C_2 - C_6 -glkenyl, C_3 - C_7 -cycloalkyl, C_3 - C_7 -cycloalkyl, and R_2 is hydrogen, with an acyl chloride of the formula R_1 -CCCI where R_1 , is C_1 - C_6 -alkyl, to form a compound of the formula I, where R_3 , R_4 , X and R_4 are as defined above and R_2 is C_1 - C_6 -alkyl-carbonyl.

h) optionally reacting a compound of the formula I, where R_b , R_b , X and n are as defined with the proviso that R_b is not hydroxy, R_1 is hydrogen, C_1 - C_6 -alkyl, C_2 - C_6 -alkenyl, C_3 - C_7 -cycloalkyl, C_3 - C_7 -cycloalkyl, wherein the phenyl group may be substituted as indicated above, and R_2 is hydrogen, with benzylchloroformate to form a compound of the formula I, where R_b , R_b , X and n are as defined R_b is as defined above and R_1 is benzyloxy carbonyl,

i) optionally reacting a compound of the formula I, where P_0 , P_4 , X and n are as defined with the proviso that P_6 is not hydroxy, P_1 is hydrogen, $C_1 - C_6$ -alikyl, $C_2 - C_6$ -alikynyl, $C_3 - C_7 - C_7$ -cycloalikyl, $C_3 - C_7 - C_7$ -cloalikenyl, phenyl- $C_1 - C_6$ -alikyl or phenyl- $C_3 - C_7 - C_7$ -cycloalikyl, wherein the phenyl group may be substituted as indicated above and P_6 is hydrogen with an isocyanate of the formula $P_1 - P_1 - C_1 - C_7 - C_$

aikyl, phenyl or phenyl- C_1 - C_6 -aikyl, wherein the phenyl group may be substituted as indicated above, to form a compound of the formula I, where R_a , R_a , X and n are as defined R_1 is as defined above and R_2 is the group

O || -C-NH-R₁₇

where R₁₇ is as defined,

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j) optionally reacting a compound of the formula I, where R_1 , R_2 , R_3 , X and n are as defined and R_4 is hydroxy with the proviso that R_2 is not C_1 - C_6 -alkylaminocarbonyl, and with a chloroformate of the formula



where R_7 is as defined to form a compound of the formula I, where R_1 , R_2 , R_3 , X and n are as defined and R_4 is the group



where R₇ is as defined,

k) optionally reacting a compound of the formula I, where R_3 , R_4 , X and n are as defined R_2 is hydrogen and R_4 is the group



where R₇ is benzyl,

with an isocyanate of the formula R_{17} -N =C=O to form a compound of the formula I, where R_1 , R_2 , R_3 , R_4 , X and I are as defined and R_2 is the group



where R_{17} is C_1 - C_6 -alkyl, phenyl or phenyl- C_1 - C_6 -alkyl, wherein the phenyl group may be substituted as indicated above,

I) optionally reacting a compound of the formula I, where R_1 , R_2 , R_3 , X and n are as defined and R_4 is hydroxy, with a compound of the formula R_1 -r-N-C-C where R_1 -r is C_1 - C_8 -alikyl, phenyl or phenyl- C_1 - C_8 -alikyl, wherein the phenyl group may be substituted as indicated in claim 1, to form a compound of the formula I, where R_1 , R_2 , R_3 , X and n are as defined and R_4 is the group



where R₁₇ is as defined above.

- A process as defined in claim 1, where n is 3.
 - 3. A compound as defined in claim 2, where
 - X is hydrogen or hydroxy.

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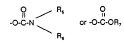
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- R₁ is hydrogen, C₃-C₆-alkynyl, C₃-C₇-cycloalkyl, phenyl or phenyl-C₁-C₆-alkyl,
- R2 is hydrogen, formyl, benzyloxycarbonyl or C1-C6-alkylaminocarbonyl,
- R₃ is hydrogen or C₁-C₆-alkyl,
- R₄ is hydrogen or a group of the formulae



wherein R_{δ} is C_1 - C_6 -alkyl or phenyl- C_1 - C_6 -alkyl and R_{δ} is hydrogen, and R_7 is phenyl- C_1 - C_6 -alkyl, where each phenyl group in the definitions of R_1 , R_{δ} and R_{δ} may be substituted as indicated in claim 1.

- 4. A process as defined in claim 3, where
 - X is hydroge
 - R₁ is C₃-C₇-cycloalkyl, phenyl-C₁-C₆-alkyl, C₃-C₆-alkynyl or phenyl-C₃-C₇-cycloalkyl, wherein the phenyl group may be substituted as indicated in claim 1,
 - Ro is hydrogen
 - R₄ is hydrogen or a group of the formula



where R₅ is C₁-C₆-alkyl and R₆ is hydrogen.

- The process as defined in claim 1, wherein 3-cyclopropylamino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]-indol-7-yl
 methylcarbonate is prepared.
- The process as defined in claim 1, wherein 4-methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent[b]-indol-7-yl methylcarbonate is prepared.

- 7. The process as defined in claim 1, wherein 1,2,3,4-tetrahydro-cyclopent(blindol-3-(2-propynyl)amine is prepared.
- 8. A process for the preparation of a compound of the formula III

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$$\mathbb{R}_{4}$$

$$\mathbb{R}_{3}$$

$$\mathbb{R}_{3}$$

$$\mathbb{R}_{n}$$

$$\mathbb{R}_{n}$$

$$\mathbb{R}_{n}$$

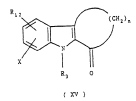
$$\mathbb{R}_{n}$$

$$\mathbb{R}_{n}$$

$$\mathbb{R}_{n}$$

where R₃, R₄, X and n are as defined in claim 1, and R₅ is hydroxy, C₁-C₈-alkoxy, amino-C₁-C₈-alkoxy, C₁-C₈-alkoy, C₃-C₇-cycloalkyl, C₃-C₇-cycloalkyl, Phenyl-C₁-G₈-alkyl, phenyl-C₃-C₇-cycloalkyl, wherein the phenyl group may be substituted as indicated in claim 1, C₁-C₈-alkylcarbonyloxy or C₁-C₈-alkylaminocarbonyloxy, or a pharmacoutically acceptable addition salt thereof, which comprises

a) reacting a compound of the formula XV



where R_3 , X and n are as defined in claim 1, and R_{12} is hydrogen, hydroxy or methoxy, with hydroxylaminhydroxhoride to form a compound of the formula III where R_3 , X and n are as defined above, R_4 has the meaning of R_2 above, and R_3 is hydroxy.

b) optionally reacting a compound of the formula III, where $R_{\rm S}$, X and n are as defined in claim 1, $R_{\rm d}$ is hydrogen, hydroxy or methoxy and $R_{\rm B}$ is hydrogen with a compound of the formula Br- $R_{\rm 13}$ -NH $_{\rm 2}$, where $R_{\rm 13}$ is $C_{\rm 1}$ - $C_{\rm S}$ -alkylene, to form a compound of the formula III where $R_{\rm 3}$, $R_{\rm 12}$, X and n are as defined and $R_{\rm B}$ is amino- $C_{\rm 1}$ - $C_{\rm G}$ -alkoxy, or

c) reacting a compound of the formula XV

where $B_{\rm g}$ X and n are as defined in claim 1 and $R_{\rm 12}$ is hydrogen, hydroxy or methoxy, with an amine of the formula NH₂ $R_{\rm 14}$ where $R_{\rm 14}$ is $C_{\rm 1}$ - $C_{\rm g}$ -alkeryl, $C_{\rm 2}$ - $C_{\rm p}$ -alkeryl, $C_{\rm 3}$ - $C_{\rm p}$ -cycloalkyol, $C_{\rm 3}$ - $C_{\rm p}$ -cycloalkyol, phenyl- $C_{\rm 3}$ - $C_{\rm p}$ -cycloalkyol or phenyl, wherein the phenyl group may be substituted as indicated in claim 1, to form a compound of the formula III, where $R_{\rm p}$ X and n are as defined above $R_{\rm q}$ has the meaning of $R_{\rm p}$ above, and $R_{\rm p}$ has the meaning of $R_{\rm p}$ above, and $R_{\rm p}$ has the meaning of $R_{\rm p}$ above,

d) optionally reacting a compound of the formula III, where R_0 , X and n are as defined in claim 1, R_4 is hydrogen, or methoxy and R_6 is hydroxy, with an isocyanate of the formula R_1 -N-C=C0 where R_1 -R1 is C_1 - C_6 -alikyl, herein the phenyl group may be substituted as indicated in claim 1, to form a compound of the formula III, where R_5 , R_4 , X and n are as defined above and R_6 is the group



where R₁₇ is as defined.

e) optionally reacting a comoound of the formula III, where R_3 , X and n are as defined in claim 1, R_4 is hydrogen, hydroxy or methoxy and R_8 is hydroxy, with an acylchloride of the formula



or an acyl anhydride of the formula $(R_1-CO)_2O$ where R_1 , $r_1 = C_1-C_6$ -alkyl, phenyl or phenyl- C_1-C_6 -alkyl, wherein the phenyl group may be substituted as indicated in claim 1, to form a compound of the formula III, where R_1 , R_2 , X and n are as defined above and R_3 is the group.



where R₁₇ is as defined above,

- 9. A process as defined in claim 8, where n is 3,
- 10. A process as defined in claim 9 where

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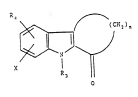
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- X is hydrogen, hydroxy or C1-C6-alkoxy
- R₃ is hydrogen or C₁-C₆-alkyl
- R4 is hydrogen, a group of the formula

- wherein R₅ is C₁-C₆-alkyl or phenyl-C₁-C₆-alkyl, R₆ is hydrogen, and R₇ is C₁-C₆-alkyl or phenyl-C₁-C₆-alkyl, R₆ is hydroxy, C₃-C₆-alkyl, amino-C₁-C₆-alkyl, C₃-C₇-cycloalkyl, phenyl-C₃-C₇-cycloalkyl, or phenyl-C₁-C₆-alkyl, where each phenyl group in the definitions of R₁, R₁ and R₂ may be substituted as indicated in claim 1.
- The process as defined in claim 8, wherein 4-methyl-3-phenylmethylimino-1,2,3,4-tetrahydrocyclopent[b]-indol-7-ol is prepared.
- Use of a compound as defined in claim 1 for the preparation of a medicament having memory dysfunction alleviating and/or antidepressant activity.
 - 13. Use of a compound as defined in claim 8 for the preparation of a medicament having antidepressant activity.
 - 14. A compound of the formula II



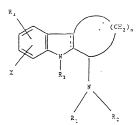
(II)

where R₃, R₄, X and n are as defined in claim 1.

50 Patentansprüche

Patentansprüche für folgende Vertragsstaaten : AT, BE, CH, DE, DK, FR, GB, IT, LI, LU, NL, PT, SE

Verbindung der Formel



in welcher

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n für 2, 3, 4 oder 5 steht;

X für Wasserstoff, C1-C6-Alkyl, C1-C6-Alkoxy, Hydroxy, Halogen, Trifluormethyl oder Nitro steht;

R, Wasserstoff, C., C_o-Aiklyl, C_o-C_o-Aikenyl, C_o-C_o-Aikenyl, Amino-C_o-C_o-Aiklyl, C_o-C_o-Cyclopalkyl, C_o-C_o-Cyclopalkyl, C_o-C_o-Cyclopalkyl, C_o-C_o-Cyclopalkyn, C_o-C_o-Cyclopalkyn, Phenyl, Phenyl-C_o-C_o-aiklyl, doder Phenyl-C_o-C_o-cyclopalkyl, wobel die Phenyigruppe mit 0, 1 oder 2 slubstituenten substituter tist, von denen jeder unabhängig voneinander C₁-C_o-Aiklyl, C₁-C_o-Aikoxy, Halogen, Trifluormethyl, Hydroxy oder Nitro ist;

ist, wobei die Gruppe "Alk" für eine bivalente C_1 - C_6 -Alkylengruppe und Y für Wasserstoff, C_1 - C_6 -Alkyl, Phenyl oder Phenyl- C_1 - C_6 -Alkyl steht, wobei die Phenylgruppe wie oben angegeben substituiert sein kann;

R₂ Wasserstoff, C₁-C₆-Alkyl, Formyl, C₁-C₆-Alkylcarbonyl, Benzyloxycarbonyl oder C₁-C₆-Alkylaminocarbonyl ist; oder aber die Gruppe



als ganzes ist

$$-N$$
 $-N$ $-N$

$$-N$$
 NH $-N$ N-C₁-C₆-Alkyl

25 oder

$$N-C_1-C_6$$
- Alkylpheny

wobei die Phenylgruppe wie oben angegeben substituiert sein kann,

$$\label{eq:prop_substitute} \begin{split} &R_3 \text{ Wasserstoff, } C_1\text{-}C_6\text{-}Alkyl, \text{ Phenyl-}C_1\text{-}C_6\text{-}alkyl, \text{ in dem die Phenylgruppe wie oben angegeben substituiert sein kann, } &C_1\text{-}C_6\text{-}Alkylcarbonyl oder } &C_1\text{-}C_6\text{-}Alkoxycarbonyl ist;} \end{split}$$

R₄ Wasserstoff, -OH,

oder

ist, wobei

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R₅ für C₁-C₆-Alkyl, C₂-C₆-Alkenyl, C₃-C₇-Cycloalkyl, C₃-C₇-Cycloalkyl, C₃-C₇-Cycloalkyl-C₁-C₆-alkyl, Phenyl, Phenyl-C₁-C₆-alkyl oder Phenyl-C₃-C₇-cycloalkyl steht, wobei die Phenylgruppe wie oben angegeben substitutiert sein kann und

 R_6 Wasserstoff, C_1 - C_6 -Alkyl, Phenyl oder Phenyl- C_1 - C_6 -Alkyl ist, wobei die Phenylgruppe wie oben angegeben substituiert sein kann; oder aber die



-Gruppe als ganzes ist

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$$-N$$

NH

 $-N$

N-C₁-C₆-Alkyl

A

 $-N$

N-C₁-C₆-Alkylphenyl

 $-N$

N-C₁-C₆-Alkylphenyl

wobei die Phenylgruppe wie oben angegeben substituiert sein kann,

Line

 R_7 C_1 - C_6 -Alkyl, Phenyl oder Phenyl- C_1 - C_6 -alkyl ist, wobei die Phenylgruppe wie oben angegeben substituiert sein kann;

unter der Bedingung, daß R_4 nicht Wasserstoff oder Hydroxy ist, wenn n für 4 oder 5 steht; oder ein pharmazeutisch verträgliches Säureadditionssalz davon.

- Verbindung gemäß Anspruch 1, in der n für 3 steht.
- 3. Verbindung gemäß Anspruch 2, in der
 - X Wasserstoff oder Hydroxy ist,
 - H₁ Wasserstoff, C₃-C₆-Alkynyl, C₃-C₇-Cycloalkyl, Phenyl oder Phenyl-C₁-C₆-alkyl ist,
 - R2 Wasserstoff, Formyl, Benzyloxycarbonyl oder C1-C6-Alkylaminocarbonyl ist,
 - R₃ Wasserstoff oder C₁-C₆-Alkyl ist,
- R₄ Wasserstoff oder eine Gruppe der Formel

$$\begin{array}{ccc} O & R_5 & O \\ -O-C-N & oder -O-C-OR_7 \end{array}$$

ist,

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in der H_S (ür C_1 - C_6 -Alkyl oder Phenyl- C_1 - C_6 -alkyl, H_6 (ür Wasserstoff und H_7 (ür Phenyl- C_1 - C_6 -alkyl steht, wobei jede Phenylgruppe in den Definitionen von H_1 , H_S und H_6 wie in Anspruch 1 angegeben substituiert sein kann.

- 4. Verbindung gemäß Anspruch 3, in der
- X Wasserstoff ist
 - R₁ C₃-C₇-Cycloalkyl, C₃-C₆-Alkynyl, Phenyl-C₃-C₇-cycloalkyl oder Phenyl-C₁-C₆-alkyl ist, wobei die Phenylgruppe wie in Anspruch 1 angegeben substituiert sein kann
 - R₂ Wasserstoff ist
 - R4 Wasserstoff oder eine Gruppe der Formel



ist

wobei R₅ für C₁-C₆-Alkyl und R₆ für Wasserstoff steht.

- Verbindung gemäß Anspruch 1, bei der es sich um 3-Cyclopropylamino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]indol-7-yl-methylcarbonat handelt.
- Verbindung gemäß Anspruch 1, die 4-Methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent[b]-indol-7-yl-methylcarbonat ist.
 - 7. Verbindung gemäß Anspruch 1, die 1,2,3,4-Tetrahydro-cyclopent[b]-indol-3-(2-propynyl)amin ist.
- 50 8. Verbindung der Formel III

$$\begin{array}{c|c} \mathbb{R}_{1} & & & & \\ \mathbb{R}_{3} & \mathbb{N} & & & \\ \mathbb{R}_{1} & & & & \\ \mathbb{R}_{1} & & & & \\ \end{array}$$

in der H_3 , H_4 , X und n die in Anspruch 1 angegebene Bedeutung zukommt und H_6 Hydroxy, C_1 - C_6 -Alkoxy, Amino- C_1 - C_6 - C_6 -Alkoxy, C_1 - C_6 -Alky, C_2 - C_6 -Alky, C_3 - C_7 - C_7 -Cycloalky, C_3 - C_7 -Cycloalky, wobei die Phenyl- C_1 - C_6 -Alky, C_1 - C_7 -Cycycloalky, wobei die Phenyl-gruppe wie in Anspruch 1 angegeben substituiert sein kann, C_1 - C_6 -Alkylota-bronyloxy oder C_1 - C_2 -Alkylaminocarbonyloxy ist, oder ein pharmazeutisch verträdliches Säureadditionssatz davon.

- 9. Verbindung gemäß Anspruch 8. in der n für 3 steht.
- 10. Verbindung gemäß Anspruch 9, in der
 - X Wasserstoff, Hydroxy oder C1-C6-Alkoxy ist
 - R₃ Wasserstoff oder C₁-C₆-Alkyl ist
 - R₄ Wasserstoff, eine Gruppe der Formel

ist.

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- in der R_5 C_1 - C_6 -Alkyl oder Phenyl- C_1 - C_6 -alkyl, R_6 Wasserstoff und R_7 C_1 - C_6 -Alkyl oder Phenyl- C_1 - C_6 -alkyl ist.
- R₈ Hydroxy, C₂-C₂-Alkylamin, Amino-C₁-C₂-alkoxy, C₁-C₂-Alkylamoptoxy, C₁-C₂-Alkylaminocarbonyloxy, C₃-C₇-Cycloalkyl, Phenyl-C₂-C₇-cycloalkyl phenyl-C₃-C₇-Cycloalkyl phenyl-C₃-C₇-Cycloalkyl phenyl-C₃-C₇-Cycloalkyl phenyl-C₃-C₇-Cycloalkyl phenyl-C₃-C₇-Cycloalkyl phenyl-C₃-Cycloalkyl phenyl-Cycloalkyl phenyl-Cyc
- Verbindung gemäß Anspruch 8, bei der es sich um 4-Methyl-3-phenylmethylimono-1,2,3,4-tetrahydrocyclopent [b]-indol-7-ol handelt.
- Pharmazeutische Zusammensetzung, die eine Verbindung gemäß Anspruch 1 oder 8 als Wirkstoff sowie eine geeignete Trägersubstanz dafür enthält.
- Anwendung einer Verbindung gemäß Anspruch 1 zur Herstellung eines Arzneimittels zur Linderung verschiedener Funktionsstörungen des Gedächtnisses und/oder mit antidepressiver Wirksamkeit.
- Anwendung einer Verbindung gemäß Anspruch 8 zur Herstellung eines Arzneimittels mit antidepressiver Wirksamkeit.
 - 15. Verfahren zur Herstellung einer Verbindung gemäß Anspruch 1, umfassend
 - a) die Reduktion einer Verbindung der Formel XVI

(XVI)

in der R₃, X und n die in Anspruch 1 angegebene Bedeutung zukommt und R₁₂ Wasserstoff, Methoxy oder Hydroxy ist, zur Bildung einer Verbindung der Formel I, in der Ra, X und n die angewiesene Bedeutung zukommt, R4 Wasserstoff, Methoxy oder Hydroxy ist, und R1 und R2 Wasserstoff sind, oder

b) die Umsetzung einer Verbindung der Formel XV

$$\begin{array}{c} \mathbb{R}_{1,2} \\ \\ \mathbb{R}_{1} \end{array} \qquad \begin{array}{c} (\mathbb{CH}_{2})_{n} \\ \\ \mathbb{R}_{3} \end{array} \qquad (XV)$$

in der R₃, X und n die in Anspruch 1 angegebene Bedeutung zukommt und R₁₂ Wasserstoff, Methoxy oder Hydroxy ist, mit Titanisopropoxid und einer Verbindung der Formel

in der die

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$$N = N$$
 No Phenyl oder $N = N$ $N = C_1 - C_6$ Alkylphenyl

wobei die Phenylgruppe wie in Anspruch 1 angegeben substituiert sein kann,

ist mit anschließender Reduktion mit Natriumborhydrid zur Bildung einer Verbindung der Formel I, in der R_3 , X und n die genannte Bedeutung haben, R_4 der für R_{12} angegebenen Bedeutung entspricht und die

als ganzes die Bedeutung der oben angegebenen

hat, oder

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c) die Reduktion einer Verbindung der Formel XVIII

in der R_0 , X und n die in Anspruch 1 angegebene Bedeutung haben, R_1 2 Wasserstoff, Methoxy oder Hydroxy ist, und R_1 4 (ür C_1 - C_6 -Alkyl, C_2 - C_6 -Alkoynl, C_3 - C_7 -Cyclobalkyl, C_7 C_7 -Cyclobalyl, C_7 -Cyclobalkyl, C_7 -Cyclobalkyl, C_7 -Cyclobalkyl, C_7

d) wahlweise die Reduktion einer Verbindung der Formel I, in der R₃, R₄, X und n die in Anspruch 1 angegebene Bedeutung haben, und R₁ und R₂ Wasserstoff sind, mit Hilfe von Boran/Tetrahydrofuran und Trifluoressigsäure zur Bildung einer Verbindung der Formel la

$$\mathbb{R}_{4}$$

$$\chi$$

$$\mathbb{R}_{3}$$

$$\mathbb{N}$$

in der Ro. Ra. X und n die angegebene Bedeutung haben.

e) wahlweise die Umsetzung einer Verbindung der Formel I, in welcher R_0 . R_4 . X und n die in Anspruch 1 angewiesene Bedeutung zukommt, und R_1 und R_2 Wasserstoff sind, mit einer Verbindung der Formel Halfl- R_1 , in der R_1 , C_1 - C_6 -Alkyl, C_2 - C_6 -Alkynyl, C_3 - C_6 -Alkynyl, C_3 - C_6 -Alkynyl, C_3 - C_6 -Alkynyl, C_3 - C_6 -Alkynyl, C_6 - C_6 -

oder

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ist, in der Alk und Y die angegebene Bedeutung haben, zur Bildung einer Gruppe der Formel I, in welcher R_0 , R_4 , X und n die bereits erwähnte Bedeutung haben, R_1 der für R_{15} oben angegebenen Bedeutung entspricht und R_5 Wasserstoff ist,

f) wahiweise die Umsetzung einer Verbindung der Formel I, in der P₃, P₄, X und n die in Anspruch 1 genannte Bedeutung haben, P₁ Wasserstoff, Cr₂-G-Mknyl, C₃-G-C-Mknyl, C₃-G-C-Cycloalkiy, C₃-G-Cy-Cycloalkiy, Phenyl-C₁-C₂-elkyl older Phenyl-C₃-G-y-cycloalkiy list, wobei die Phenylgruppe wie in Anspruch 1 angegeben substituiert sein kann, und P₂ Wasserstoff ist, mit Ameisensäure zur Bildung einer Verbindung der Formel I, in welcher P₃, P₄, X und n die in Anspruch 1 angewiesene Bedeutung zukommt, P₁ die oben genannte Bedeutung hat und R₃-Formyl ist.

g) wahlweise die Umsetzung einer Verbindung der Formel I, in der R₃, R₄, X und n die in Anspruch 1 genannte Bedeutung haben, R₁ Wasserstoff, C₁-C₅-Alkyl, C₂-C₅-Alkynyl, C₃-C₇-Cyclotalkyl, St. wobei die Phenylgruppe wis in Anspruch 1 angegeben substituiert sein kann, und R₂ Wasserstoff ist, mit einem Acylchlorid der Formel R₁₇-COCi, in der R₁₇-C₇-C₈-Alkyl ist, zur Bildung einer Vorbindung der Formel I, in der R₃, R₄. X und n die in Anspruch 1 angewiesene Bedeutung aktuent, R₁-C₁-C₈-Alkyl(vachoryl ist, zur Cyclotalkyl).

h) wahlweise die Umsetzung einer Verbindung der Formel I, in der R₃, R₄, X und n die in Anspruch 1 genannte Bedeutung haben, unter der Voraussetzung, daß R₄ nicht Hydroxy, R₁, Wasserstoft, C₁-C₆-Alikly, C₂-C₆-Cycloalikly, C₂-Cy-Cycloalikly, C₈-Cy-Cycloalikly, C₈-Cy-Cycloalikly, der Phenyl-Cy-C₇-Cycloalikly, ist, wobei die Phenylgruppe wie in Anspruch 1 angegeben substituiert sein kann, und R₂ Wasserstoff ist, mit einem Benzylchloroformat zur Bildung einer Verbindung der Formel I, in der R₃, R₄, X und n die oben angegebene Bedeutung haben, R, die oben genannte Bedeutung hab tun dR, Benzyloxycarabonyl ist, die oben genannte Bedeutung hab tun dR, Benzyloxycarabonyl ist.

i) wahlweise die Umsetzung einer Verbindung der Formel I, in der R₃, R₄, X und n die in Anspruch 1 genannte Bedeutung haben, unter der Voraussetzung, daß R₄ nicht Hydroxy, R₁ Wasserstoft, C₇-C₈-Alklyi, C₂-C₆-Alkeynyl, C₃-C₇-cycloalkyl, C₃-C₇-Cycloalkyl, C₃-C₇-Cycloalkyl und R₇-Wasserstoff ist, mit Isocyanat der Formel R₇₇-Ne-C₉-C₁, dem R₇₇-C₇-C₆-Alklyl, Phenyl oder Phenyl-C₃-C₇-Cycloalkyl

 C_1 - C_2 -alkyl ist, wobei die Phenylgruppe in der Definition von R_1 und $R_{1,7}$ wie in Anspruch 1 angegeben substitutien sein kann, zur Bildung einer Vorbindung der Formel I, in der R_3 , R_4 , X und n die oben angegebene Bedeutung haben, R_1 , die oben genannte Bedeutung hat und R_2 die

ist, in der R₁₇ die angewiesene Bedeutung zukommt,

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j) wahlweise die Umsetzung einer Verbindung der Formel I, in der R_1 , R_2 , R_3 , X und n die in Anspruch 1 genannte Bodeutung haben und R_1 Hydroxy ist, unter der Voraussetzung, GaB R_2 nicht C_1 - C_6 -Alkylaminocarbonyl ist, mit einem Chloroformat der Formel

in der R_7 die in Anspruch 1 genannte Bedeutung hat, zur Bildung einer Verbindung der Formel I, in der R_1 , R_2 , R_3 , X und n die oben angegebene Bedeutung haben und R_4 die

ist, in der R7 die in Anspruch 1 angewiesene Bedeutung zukommt,

k) wahlweise die Umsetzung einer Verbindung der Formel I, in der R_1 , R_3 , X und n die in Anspruch 1 genannte Bedeutung haben, R_2 Wasserstoff und R_4 die

ist, in der R_7 Benzyl ist, mit Isocyanat der Formel R_{17} -N=C=O, zur Bildung einer Verbindung der Formel I, in der R_1 , R_2 , R_3 , R_4 , X und n die oben angegebene Bedeutung haben und R_2 die

ist, in der R₁₇ C₁-C₈-Alkyl, Phenyl oder Phenyl-C₁-C₆-alkyl ist, wobei die Phenylgruppe wie in Anspruch 1 angegeben substitutiert sein kann,

I) wahlweise die Umsetzung einer Verbindung der Formel I, in der R₁, R₂, R₃, X und n die in Anspruch 1 genannte Bedeutung haben und R₄ Hydroxy ist, mit einer Verbindung der Formel R₁₇-N-C=-N in der R₁₇ C₁-C₃-C₃-Alkyl, Phenyl oder Phenyl-C₁-C₃-alkyl ist, wobel die Phenylgruppe wie in Anspruch 1 angegeben substitutiert sein kann, zur Bildung einer Verbindung der Formel I, in der R₁, R₂, R₃, X und n die oben angegebene Bedeutung haben und R₂, die

O || -O-C-NH-R,,-Gruppe

ist, in der R₁₇ die obengenannte Bedeutung zukommt.

16. Verfahren zur Herstellung einer Verbindung der Formel III gemäß Anspruch 8. umfassend

a) die Umsetzung einer Verbindung der Formel XV

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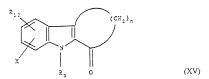
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in der H_3 . X und n die in Anspruch 1 angegebene Bedeutung zukommt und H_{12} Wasserstoff, Hydroxy oder Methoxy ist, mit Hydroxylaminhydrochlorid zur Bildung einer Verbindung der Formel III, in der H_3 . X und n die oben angewiesene Bedeutung zukommt, H_4 der für H_{12} oben angegebenen Bedeutung enispricht und H_3 Hydroxy ist,

b) wahlweise die Umsetzung einer Verbindung der Formel III, in welcher R₃, X und n die in Anspruch 1 angewiesene Bedeutung zukommt, R₆ Wasserstoff, Hydroxy oder Methoxy und R₆ Wasserstoff ist, mit einer Verbindung der Formel Br-R₁₋₃NH₂, in der R₁₋₃ (fir C₁-C₆-Alkylen steht, zur Bildung einer Verbindung der Formel IIII, in der R₈, H₁₂, X und n die genannte Bedeutung haben und R₆ Aminc-C₁-C₆-alkoxy ist, oder

c) die Umsetzung einer Verbindung der Formel XV



in dor Π_{2} , X und n die in Anspruch 1 angegebene Bedeutung zukommt und Π_{12} Wasserstoff, Methoxy oder Hydroxy ist, mit einem Amin der Formel NH₂R₁₄, in der Π_{14} Cr₂-C₅-Alkyl, C₂-C₆-Alkeyl, C₃-C₇-Cycloalkyn, C₃-Cycloalkyn, C₃-Cy

d) wahlwaise die Umsetzung einer Verbindung der Formel III, in welcher R_0 , X und n die in Anspruch 1 angewiesene Bedeutung zukommt, R_4 Wasserstolf oder Methoxy und R_6 Hydroxy ist, mit Isocyanat der Formel R_{17} -N-C- Ω , in der R_{17} - Ω - Ω - Ω -Miyl, Phenyl oder Phenyl- Ω - Ω - Ω -althyl ist, wobei die Phenylgruppe wie in An-

spruch 1 angegeben substitutiert sein kann, zur Bildung einer Verbindung der Formel III, in der R_3 , R_4 , X und n die oben genannte Bedeutung haben und R_8 die

O II R.,-NH-C-O-Gruppe

ist, in der R₁₇ die angegebene Bedeutung hat,

 e) wahlweise die Umsetzung einer Verbindung der Formel III, in der R₃. X und n die in Anspruch 1 genannte Bedeutung haben, R₄ Wasserstoff, Hydroxy oder Methoxy und R₆ Hydroxy ist, mit einem Acylchlorid der Formel

> O || R₁₇-C-Cl

oder einem Säureanhydrid der Formel (R_{17} CO)₂O, in denen R_{17} für C_{1} - C_{6} -Alkyl, Phenyl oder Phenyl- C_{17} - C_{6} -alkyl steht, wobei die Phenylgruppe wie in Anspruch 1 angegeben substituiert sein kann, zur Bildung einer Verbindung der Formel III, in der R_{18} , R_{14} X und n die oben angegebene Bedeutung haben und R_{15} die Axionium der Formel III, in der R_{15} , R_{14} X und n die oben angegebene Bedeutung haben und R_{15} die

(II)

ist, in der R₁₂ die oben genannte Bedeutung hat,

17. Verbindung der Formel II

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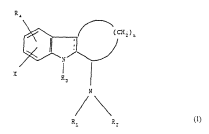
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in der R₃, R₄, X und n die in Anspruch 1 genannte Bedeutung haben.

Patentansprüche für folgende Vertragsstaaten : ES, GR

1. Verfahren zur Herstellung einer Verbindung der Formel I



in welcher

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n für 2, 3, 4 oder 5 steht;

X für Wasserstoff, C1-C6-Alkyl, C1-C6-Alkoxy, Hydroxy, Halogen, Trifluormethyl oder Nitro steht;

 F_{ij} Wasserstoff, C_{ij} - C_{ij} -Alkyl, C_{ij} - C_{ij} -Alkyl, F_{ij} - F_{ij} F_{ij} -

ist, wobei die Gruppe *Alk' für eine bivalente C_1 - C_6 -Alkylengruppe und Y für Wasserstoff, C_1 - C_6 -Alkyl, Phenyl oder Phenyl- C_1 - C_6 -alkyl steht, wobei die Phenylgruppe wie oben angegeben substituiert sein kann;

 R_2 Wasserstoff, $C_1^-C_6$ -Alkyl, Formyl, $C_1^-C_6^-$ Alkyl, Earbonyl, Benzyloxycarbonyl oder $C_1^-C_6^-$ Alkylaminocarbonyl ist; oder aber die Gruppe



55 als ganzes ist

$$N$$
 N N

oder

wobei die Phenylgruppe wie oben angegeben substituiert sein kann,

R₃ Wasserstoff, C₁-C₆-Alkyl, Phenyl-C₁-C₆-alkyl, in dem die Phenylgruppe wie oben an gegeben substituiert sein kann, C₁-C₆-Alkylcarbonyl oder C₁-C₆-Alkoxycarbonyl ist; R4 Wasserstoff, -OH,

oder

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_0.c.o

ist, wobei

 R_g^- für C_q - C_g -Alkyl, C_2 - C_g -Alkenyl, C_3 - C_g -Alkynyl, C_3 - C_7 -Cycloalkyl, C_3 - C_7 -Cycloalkyl- C_1 - C_g -alkyl, Phenyl- C_3 - C_7 -cycloalkyl steht, wobel die Phenylgruppe wie oben angegeben substitutiert sein kann, und

 H_6 Wasserstoff, C_1 - C_6 -Alkyl, Phenyl oder Phenyl- C_1 - C_6 -alkyl ist, wobei die Phenylgruppe wie oben angegeben substituiert sein kann;

oder aber die

$$-N$$

-Gruppe als ganzes ist

$$-N$$
 $-N$ $-N$ 0

$$-N \qquad NH \qquad -N \qquad N-C_1 \cdot C_6 \cdot A 1 ky 1$$

wobei die Phenylgruppe wie oben angegeben substituiert sein kann,

und

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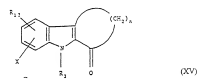
 R_7C_1 - C_6 -Alkyl, Phenyl oder Phenyl- C_1 - C_6 -alkyl ist, wobei die Phenylgruppe wie oben angegeben substituiert sein kann:

- unter der Bedingung, daß R₄ nicht Wasserstoff oder Hydroxy ist, wenn n für 4 oder 5 steht; oder eines pharmazeutisch verträglichen Säureadditionssalzes davon, umfassend
 - a) die Reduktion einer Verbindung der Formel XVI

in der H_3 . X und n die in Anspruch 1 angegebene Bedeutung zukommt und H_{12} Wasserstoff, Meihoxy oder Hydroxy ist, zur Bildung einer Verbindung der Formel I. in der H_3 . X und n die angewiesene Bedeutung zukommt, H_3 Wasserstoff, Methoxy oder Hydroxy ist, und H_3 und H_3 Wasserstoff sind, oder

(XVI)

b) die Umsetzung einer Verbindung der Formel XV



in der R₃, X und n die in Anspruch 1 angegebene Bedeutung zukommt und R₁₂ Wasserstoff, Methoxy oder Hydroxy ist, mit Titanisopropoxid und einer Verbindung der Formel

50 in der die

N A-Gruppe



$$-N$$
 NH $-N$ $N-C_1-C_{6^-}$ Alkyl

20 wobei die Phenylgruppe wie in Anspruch 1 angegeben substituiert sein kann,

ist, mit anschließender Reduktion mit Natriumborhydrid zur Bildung einer Verbindung der Formel I, in der $R_{\rm gr}$ X und n die genannte Bedeutung haben, $R_{\rm 4}$ der für $R_{\rm 12}$ angegebenen Bedeutung entspricht und die

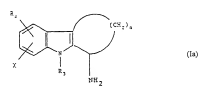
als ganzes die Bedeutung der oben angegebenen

hat, oder

c) die Reduktion einer Verbindung der Formel XVIII

in der R_b X und n die in Anspruch 1 engegebene Bedeutung haben, R_{12} Wasserstoff, Methoxy oder Hydroxy ist, und R_{14} für C_1 - C_6 -Alkyl, C_2 - C_6 -Alkonyl, C_1 - C_6 -Alkynyl, C_3 - C_7 -Cycloalkyl, C_3 - C_7 - C_7 -Cycloalkyl, C_3 - C_7

d) wahlweise die Reduktion einer Verbindung der Formel I, in der R₃. R₄. X und n die in Anspruch 1 angegebene Bedeutung haben, und R₁ und R₅ Wasserstoff eind, mit Hilfe von Boran/Totrahydrofuran und Trifluoressigsäure zur Bildung einer Verbindung der Formel II.



in der R₃, R₄, X und n die angegebene Bedeutung haben,

e) wahlweise die Umsetzung einer Verbindung der Formel I, in weicher R_3 , R_4 , X und n die in Anspruch 1 angewiesene Bedeutung zukommt, und R_1 und R_2 Wasserstoff sind, mit einer Verbindung der Formel Half- R_5 in der R_1 , C_1 - C_6 -Alkyl, C_2 - C_6 -Alkyl, C_2 - C_6 -Alkyl, C_2 - C_6 -Alkyl, C_7 - C_7 -Oser (C_7 C_7 - C_7 - C_7 - C_7 -Oser (C_7 - C_7 - C_7 - C_7 -Oser (C_7 - C_7 - C_7 - C_7 -Oser (C_7 - C_7 - C_7 - C_7 -Oser (C_7 - C_7 - C_7 - C_7 -Oser (C_7 - C_7 - C_7 - C_7 - C_7 -Oser (C_7 - C_7 - C_7 - C_7 -Oser (C_7 - C_7 - C_7 - C_7 - C_7 -Oser (C_7 - C_7 - C_7 - C_7 -Oser (C_7 - C_7 - C_7 - C_7 - C_7 - C_7 -Oser (C_7 - $C_$

$$-Alk-N$$
, $-Alk-N$, $-Alk-N$, $-Alk-N$,

oder

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ist, in der Alk und Y die angegebene Bedeutung haben, zur Bildung einer Gruppe der Formel I, in welcher P_0 . P_4 , X und n die bereits erwähnte Bedeutung haben, P_1 der für P_1 5 oben angegebenen Bedeutung entspricht und P_3 Wasserstoff ist.

 $\label{eq:continuous} \begin{tabular}{l} (1) washiweise die Umsetzung einer Verbindung der Formel II, in der <math>B_0$, B_4 , X und n die in Anspruch 1 genannte Bedeutung haben, B_1 Wasserstoff, C_1 - C_0 -Alkiyl, C_2 - C_0 -Alkiynyl, C_3 - C_0 -C-ycc)-clocklayl, B_1 -Alkiynyl, C_3 - C_0 -ycc)-clocklayl, B-Aly, who die B-Aly B

g) wahlweise die Umsetzung einer Verbindung der Formel I, in der R₃, R₄, X und n die genannte Bedeutung haben, R₁ Wasserstoff, C₃-C₅-Alkyl, C₂-C₅-Alkonyl, C₃-C₅-Alkynyl, C₃-C₅-Cycloalkyl, Phenyl-C₄-C₅-q-Qistellkyl jut, Wobei die Phenylgrupe wie in Anspruch 1 angegeben substitutiert sein kann, und R₂ Wasserstoff ist, mit einem Acylchlorid der Formel R₁₇-COI, in der R₁₇ C₁-C₆-Alkyl ist, zur Bildung einer Werbindung der Formel I, in der R₃, R₄, X und n die in Anspruch 1 angewiesene Bedeutung auch die S, C₁-C₆-Alkyl ist, var Bildung einer Werbindung der Formel I, in der R₃, R₄, X und n die in Anspruch 1 angewiesene Bedeutung auch B. C₁-C₆-Alkyl ist, Alkyl ist, var Bildung einer Werbindung der Formel I, in der R₃, R₄, X und n die in Anspruch 1 angewiesene

h) wahlweise die Umsetzung einer Verbindung der Formel I, in der R_3 , R_4 , X und n die genannte Bedeutung haben, unter der Voraussetzung, das R_3 nicht Hydroxy, R_1 , Wasserstollt, $C_1 \cdot C_2 \cdot C_3 \cdot Alkenyl, C_2 \cdot C_3 \cdot C_4 \cdot C_6 \cdot Alkenyl, C_3 \cdot C_7 \cdot C_7 \cdot C_6 \cdot Alkenyl, C_3 \cdot C_7 \cdot C_7 \cdot C_6 \cdot Alkenyl, C_3 \cdot C_7 \cdot C_7 \cdot C_6 \cdot Alkenyl, C_3 \cdot C_7 \cdot C$

i) wahiweisa die Umsatzung einer Verbindung der Formel I, in der R_0 , R_0 , X und n die genannte Bedoutung haben, unter der Voraussetzung, daß R_1 nicht Hydroxy, R_1 Wassersstoff, C_2 - C_3 -Rilky, C_2 - C_2 -Alley, R_1 - R_2 -Rilky, R_2 - R_2

ist, in der R₁₇ die angewiesene Bedeutung zukommt,

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j) wahlweise die Umsetzung einer Verbindung der Formel I, in der R_1 , R_2 , R_3 , X und n die genannte Bedeutung haben und R_4 Hydroxy ist, unter der Voraussetzung, daß R_2 nicht C_1 - C_6 -Alkylaminocarbonyl ist, mit einem Chloroformat der Formel

in der R₇ die genannte Bedeutung hat, zur Bildung einer Verbindung der Formel I, in der R₁, R₂, R₃. X und n die oben angegebene Bedeutung haben und R₄ die

ist, in der R7 die angewiesene Bedeutung zukommt,

k) wahlweise die Umsetzung einer Verbindung der Formel I, in der R_1 , R_3 , X und n die genannte Bedeutung haben, R_2 Wasserstoff und R_4 die

O ll -O-C-OR₂-Gruppe

ist, in der R7 Benzyl ist, mit Isocyanat der Formel R17-N=C=O,

zur Bildung einer Verbindung der Formel I, in der R_1 , R_2 , R_3 , R_4 , X und n die angegebene Bedeutung haben und R_5 die

ist, in der R₁₇ C₁-C₆-Alkyl, Phenyl oder Phenyl-C₁-C₆-alkyl ist, wobei die Phenylgruppe wie oben angegeben substituiert sein kann,

i) wahlweise die Umsetzung einer Verbindung der Formel I, in der R_1 , R_2 , R_3 , X und n die genannte Bedeutung haben und R_4 Hydroxy ist, mit einer Verbindung der Formel R_1 - \mathbb{P}_1 - \mathbb{P}_2 - \mathbb{P}_3 - \mathbb{P}_3 , \mathbb{P}_3 - \mathbb{P}_4 - \mathbb{P}_3 - \mathbb{P}_4 - \mathbb



ist, in der R₁₇ die obengenannte Bedeutung zukommt.

- Verfahren gemäß Anspruch 1, in dem n für 3 steht.
- 35 3. Verfahren gemäß Anspruch 2. in der

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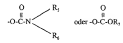
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- X Wasserstoff oder Hydroxy ist,
- R₁ Wasserstoff, C₃-C₆-Alkynyl, C₃-C₇-Cycloalkyl, Phenyl oder Phenyl-C₁-C₆-alkyl ist,
- R₂ Wasserstoff, Formyl, Benzyloxycarbonyl oder C₁-C₆-Alkylaminocarbonyl ist,
 - R₃ Wasserstoff oder C₁-C₆-Alkyl ist,
 - R₄ Wasserstoff oder eine Gruppe der Formel



ist

in der R₅ für C₁-C₆-Alkyl oder Phenyl-C₁-C₆-alkyl, R₆ für Wasserstoff und R₇ für Phenyl-C₁-C₆-alkyl steht, wobei jede Phenylgruppe in den Definitionen von R₁, R₅ und R₆ wie in Anspruch 1 angegeben substituiert sein kann.

- 4. Verfahren gemäß Anspruch 3, in dem
 - X Wasserstoff ist

- R₁ C₃-C₇-Cycloalkyl, Phenyl-C₁-C₆-alkyl, C₃-C₆-Alkynyl oder Phenyl-C₃-C₇-cycloalkyl ist, wobei die Phenylgruppe wie in Anspruch 1 angegeben substituiert sein kann
- R₂ Wasserstoff ist

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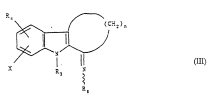
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R₄ Wasserstoff oder eine Gruppe der Formel



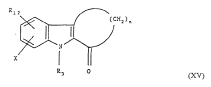
ist, wobei R₅ für C₁-C₆-Alkyl und R₅ für Wasserstoff steht.

- Verfahren gemäß Anspruch 1, bei dem 3-Cyclopropylamino-4-methyl-1,2,3,4-tetrahydrocyclopent[b]-indoi-7-yi-methylcarbonat heroestellt wird.
- Verfahren gemäß Anspruch 1, bei dem 4-Methyl-3-phenylmethylamino-1,2,3,4-tetrahydrocyclopent[b]-indol-7-ylmethylcarbonat hergestellt wird.
 - 7. Verfahren gemäß Anspruch 1, bei dem 1.2,3,4-Tetrahydro-cyclopent[b]-indol-3-(2-propynyt)amin hergestellt wird.
 - 8. Verfahren zur Herstellung einer Verbindung der Formel III



in dem R_5 , R_4 , X und n die in Anspruch 1 angegebene Bedeutung zukommt und R_6 Hydroxy, C_1 - C_6 -Alkoy, Amino- C_1 - C_6 -Alkoy, C_3 - C_6 -Cycloalkyl, C_3 - C_7 -Cycloalkyl, C_3 - C_7 -Cycloalkyl, Phenyl- C_1 - C_7 -alkyl oder Phenylgruppe wie in Anspruch 1 angegeben substitutiert sein kann, C_1 - C_6 -Alkylcarbonyloxy oder C_1 - C_6 -Alkylaminocarbonyloxy ist, oder eines pharmazeutisch verträglichen Säureaddfilonssatzes davon, umfassend

a) die Umsetzung einer Verbindung der Formel XV



in der H_3 , X und n die in Anspruch 1 angegebene Bedeutung zukommt und H_{12} Wasserstoff, Hydroxy oder Melhoxy ist, mit Hydroxylaminhydrochlorid zur Bildung einer Verbindung der Formel III, in der H_3 , X und n die oben angewiesene Bedeutung zukommt, H_4 der für H_{12} oben angegebenen Bedeutung enlspricht und H_3 Hydroxy ist.

b) wahlweise die Umsetzung einer Verbindung der Formel III, in welcher R₃, X und n die in Anspruch 1 angewiesene Bedeutung zukornmt, R₄ Wasserstoff, Hydroxy oder Methoxy und R₅ Wasserstoff ist, mit einer Verbindung der Formel Br-R₁₃-NH₂, in der R₁₃ (für C₇-C₆-Alkylen steht, zur Bildung einer Verbindung der Formel IIII, in der R₅, H₁₂, X und n die genannte Bedeutung haben und R₆ Amino-C₇-C₆-alkoxy ist, oder

c) die Umsetzung einer Verbindung der Formel XV

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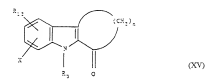
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in der R_0 , X und n die in Anspruch 1 angegebene Bedeutung zukommt und R_{12} Wasserstoff, Melhoxy oder Hydroxy lst, mit einem Anrin der Formel NH₂R₁₄, in der R_{14} C₁-Ce₃-Alkyl, C₂-Ce₄-Alkenyl, C₃-Ce₅-Alkynyl, C₃-Ce₇-Cycloalkyl, Ce₃-Ce₇-Cycloalkyl, Phenyl-C₁-Ce₃-alkyl, Phenyl-C₃-Ce₇-Cycloalkyl oder Phenyl ist, wobei die Phenylgruppe wie in Anspruch 1 angegeben substituiert sein kann, zur Bildung einer Verbindung der Formel III, in der R_3 , X und n die oben genannte Bedeutung haben, R_4 die Bedeutung von R_{12} oben hat und R_6 für R_{14} , oben steht,

d) wahlweise die Umsetzung einer Verbindung der Formel III, in welcher $R_{\rm h}$, X und n die in Anspruch 1 angewiesene Bedeutung zukommt, $R_{\rm d}$ Wassertsolf oder Methoxy und $R_{\rm B}$ Hydroxy ist, mit Isocyanat der Formel $R_{\rm 17}$ -N-C-C-D, in der $R_{\rm T}$ Or- $Q_{\rm e}$ -Alkyl, Phornyl oder Phornyl-C- $Q_{\rm e}$ -Alkyl ist, woboi die Pherrylgruppe wie in Anspruch 1 angegeben substituient sein kann, zur Bildung einer Verbindung der Formel III, in der $R_{\rm 3}$, $R_{\rm d}$, X und n die oben genannte Bedeutung haben und $R_{\rm d}$ die

ist, in der R₁₇ die angegebene Bedeutung hat,

 e) wahlweise die Umsetzung einer Verbindung der Formel III, in der R₃. X und n die in Anspruch 1 genannte Bedeutung haben, R₄ Wasserstoff, Hydroxy oder Methoxy und R₈ Hydroxy ist, mit einem Acylchlorid der Formel

oder einem Saureanhydrid der Formel $(R_{17}\text{-CO})_2\text{O}$, in denen R_{17} für C_1 - C_8 -Alkyl, Phenyl oder Phenyl- C_1 - C_6 -alkyl seht, wobei die Phenylgruppe wie in Anspruch 1 angegeben substitutier sein kann, zur Bildung einer Verbindung der Formel III, in der R_3 - R_4 . X und n die oben angegebene Bedeutung haben und R_6 die

ist, in der R₁₇ die oben genannte Bedeutung hat,

- 9. Verfahren gemäß Anspruch 8. in dem n für 3 steht.
- 10 10. Verfahren gemäß Anspruch 9, in dem
 - X Wasserstoff, Hydroxy oder C1-C6-Alkoxy ist
 - R₃ Wasserstoff oder C₁-C₆-Alkyl ist
 - R4 Wasserstoff, eine Gruppe der Formel

ist

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- in der R₅ C₇-C₆-Alkyl oder Phenyl-C₁-C₆-alkyl, R₆ Wasserstoff und R₇ C₁-C₆-Alkyl oder Phenyl-C₁-C₆-alkyl ist.

 R₆ Hydroxy, C₂-C₆-Alkyl, Arino-C₁-C₆-alkyx, C₃-C₆-Alkylearbonyloxy, C₁-C₆-Alkylaminocarbonyloxy, C₂-C₇-Cycloalkyl, Phenyl-C₂-C₇-cycloalkyl, Phenyl-C₃-C₇-cycloalkyl, Phenyl-
- Verlahren gemäß Anspruch 8, bei dem 4-Methyl-3-phenylmethylimono-1,2,3,4-tetrahydrocyclopent[b]-indol-7-ol
 hergestellt wird.
 - Anwendung einer Verbindung gemäß Anspruch 1 zur Herstellung eines Arzneimittels zur Linderung verschiedener Funktionsstörungen des Gedächtnisses und/oder mit antidepressiver Wirksamkeit.
- 35 13. Anwendung einer Verbindung gemäß Anspruch 8 zur Herstellung eines Arzneimittels mit antidepressiver Wirksamkeit
 - 14. Verbindung der Formel II

R₄ (CH₂

(II)

in der R₃, R₄, X und n die in Anspruch 1 genannte Bedeutung haben.

Revendications

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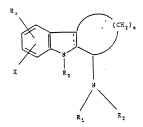
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Revendications pour les Etats contractants suivants : AT, BE, CH, DE, DK, FR, GB, IT, LI, LU, NL, PT, SE

Composé de formule



dans laquelle

- n est 2, 3, 4 ou 5;
- X est un atome d'hydrogène ou d'halogène ou un groupe alkyle en C₁-C₆, alcoxy en C₁-C₆, hydroxy, trifluoro-méthyle ou nitro;
- Fig. ast un atome d'hydrogène ou un groupe aiktyle en C₁-C₆, alcényle en C₂-C₆, alcynyle en C₃-C₆, aminoaiktyle en C₃-C₆, alcynyle en C₃-C₆, alcynyle en C₃-C₆, aminoaiktyle (C₁-C₆), alcynication (C₃-C₇), elsyne (C₁-C₆), cycloaiktyle (C₁-C₆), cycloaiktyle (C₁-C₆), cycloaiktyle en C₃-C₇, phányle, phányl-alktyle (C₁-C₆) ou phányl-cycloaiktyle (C₃-C₇), le groupe phányle distant substitué par 0, 1 ou 2 substituants, représentant chacun indépendamment un atome d'halogène ou un groupe alktyle en C₁-C₆, alcoxy en C₁-C₆, irflutorométhyle, hydroxy ou nife.

"alk" représentant un groupe alkylène divalent en $C_1 \cdot C_0$, et Y représentant un alome d'hydrogène ou un groupe alkyle en $C_1 \cdot C_0$, phényle ou phényl-alkyle $(C_1 \cdot C_0)$, le groupe phényle pouvant être substitué comme indiqué plus haut;

R₂ est un alcome d'hydrogène ou un groupe alkyle en C₁-C₆, formyle, alkyl(C₁-C₆)-carbonyle, benzyloxycarbonyle ou alkyl(C₁-C₆)-amino-carbonyle, ou bien le groupe

dans son ensemble est

$$-N$$
 NH $-N$ N-C₁-C₆-alkyle

le radical phényle pouvant être substitué comme indiqué plus haut,

- $\label{eq:proposed_equation} \begin{array}{ll} \mathsf{Fl}_3 & \text{est un atome d'hydrogène ou un groupe alkyle en } C_1^-C_6, \text{phényl-alkyle}(C_1^-C_6), \text{le fragment phényle pouvant être substitué comme indiqué plus haut, alkyl(} C_1^-C_6)-\text{carbonyle ou alcoxy}(C_1^-C_6)-\text{carbonyle}; \end{array}$
- R₄ est un atome d'hydrogène, -OH, un groupe

ou

dans lequel

- H_5 est un radical alkyle en $C_1 \cdot C_6$, alcényle en $C_2 \cdot C_6$, alcynyle en $C_3 \cdot C_6$, cycloalkyle en $C_3 \cdot C_7$, cycloalkyle $(C_3 \cdot C_7)$ -alkyle $(C_1 \cdot C_6)$, by hényle-gycloalkyle $(C_3 \cdot C_7)$, le fragment phényle pouvant for s eubstitué comme indiqué blus haut; et
- Fig. est un atome d'hydrogène ou un groupe alkyle en C₁-C_e, phényle ou phényl-alkyle(C₁-C_e), le fragment phényle pouvant être substitué comme indiqué plus haut; ou bien le groupe

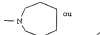
est dans son ensemble

$$-\sqrt{-\lambda}$$

$$N - N$$
 NH $N - N$ $N - C_1 - C_6$ -alkyle

$$- \text{ N-phényle} \quad \text{ou} \quad \\ N-C_1 \cdot C_{5^-} \text{ alkylphényle} \, .$$

le fragment phényle pouvant être substitué comme indiqué plus haut,



.

B₇ est un groupe alkyle en C₁-C₆, phényle ou phényl-alkyle(C₁-C₆), le fragment phényle pouvant être substitué comme indiqué plus haut:

étant entendu que R₄ n'est pas un atome d'hydrogène ou le groupe hydroxy lorsque n est 4 ou 5; ou sel d'addition avec un acide pharmaceutiquement acceptable de celui-ci.

- Composé selon la revendication 1, dans lequel n est égal à 3.
 - 3. Composé selon la revendication 2, dans lequel
 - X est un atome d'hydrogène ou le groupe hydroxy.
 - R₁ est un atome d'hydrogène ou un groupe alcynyle en C₃-C₆, cycloalkyle en C₃-C₇, phényle ou phényl-alkyle
 - R2 est un atome d'hydrogène ou un groupe formyle, benzyloxycarbonyle ou alkyl(C1-Ca)amino-carbonyle,
 - R₃ est un atome d'hydrogène ou un groupe alkyle en C₁-C₆,
 - R4 est un atome d'hydrogène ou un groupe de formule

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formules dans lesquelles R_e est un groupe alkyle en C_1 - C_6 ou phényl-alkyle(C_1 - C_6) et R_6 est un atome d'hydrogène. et R_1 est un groupe phényl-alkyle(C_1 - C_6), chaque groupe phényle dans les définitions de R_1 , R_5 et R_6 pouvant être substitué comme indiqué dans la revendication 1.

- 4. Composé selon la revendication 3, dans leguel
 - X est un atome d'hydrogène,
 - R₁ est un groupe cycloalikyle en C₃·C₇, alcynyle en C₃·C₆, phényl-cycloalkyle(C₃·C₇) ou phényl-alkyle(C₁·C₆), le fragment phényle pouvant être substitué comme indiqué dans la revendication 1,
 - Ro est un atome d'hydrogène,
 - R. est un atome d'hydrogène ou un groupe de formule

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- dans laquelle R₅ est un groupe alkyle en C₁-C₆ et R₆ est un atome d'hydrogène.
- Composé selon la revendication 1, qui est le méthylcarbonate de 3-cyclopropylamino-4-méthyl-1, 2 3,4-tétrahydrocyclopent[b]indole-7-yle.
- Composé selon la revendication 1, qui est le méthylcarbonate de 4-méthyl-3-phénylméthylamino-1, 2 3,4-tétrahydrocyclopent/folindole-7-yle.
 - 7. Composé selon la revendication 1, qui est la 1,2,3,4-tétrahydrocyclopent[b]indole-3-(2-propynyl)amine.

8. Composé de formule III

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(CH₂)_n
(X)

R₃

R₄

dans iaquelle P_0 , P_0 , X et n sont tels que définis dans la revendication 1, et P_0 est un groupe hydroxy, alcoxy en C_1 - C_0 , arminoalcoxy(C_1 - C_0), alkyle en C_1 - C_0 , alcynyle en C_3 - C_0 , cycloalkyle en C_3 - C_0 , cycloalkyle en C_3 - C_0 , cycloalkyle en C_3 - C_0 , opheryl-alkyle en C_3 - C_0 - C_0), to fragment phényle provant être substitué comme indiqué dans la revendication 1, alkyle(C_1 - C_0 -earbonyloxy ou alkyl(C_1 - C_0 -aminocarbonyloxy, ou sel d'addition pharma-ceutiquement acceptable de cellui-ci.

- 9. Composé selon la revendication 8, dans lequel n est égal à 3.
- 25 10. Composé selon la revendication 9, dans leguel
 - X est un atome d'hydrogène ou un groupe hydroxy ou alcoxy en C₁-C₆.
 - R₃ est un atome d'hydrogène ou un groupe alkyle en C₁-C₆,
 - R4 est un atome d'hydrogène ou un groupe de formule

formules dans lesquelles R_5 est un groupe alkyle en C_1 - C_6 ou phényl-alkyle(C_1 - C_6), R_6 est un atome d'hydrogène. et R_7 est un groupe alkyle en C_1 - C_6 ou phényl-alkyle(C_1 - C_6),

- R₈ est un groupe hydroxy, alcynyle en C₃·C₆, amino-alcoxy(C₁·C₆), alkyl(C₁·C₆)-carbonyloxy, alkyl(C₁·C₆)aminocarbonyloxy, cycloalkyle en C₃·C₇, phényl-cycloalkyle(C₂·C₇) ou phényl-alkyle(C₁·C₆), chaque fragment phényle dans le selémitions de R₂, E₆ in R₆ pouvant être substitute comme indicuel dans la revendication 1.
- Composé selon la revendication 8, qui est le 4-méthyl-3-phénylméthylimino-1,2,3,4-tétrahydrocyclopent[b]indole-7-ol.
- Composition pharmaceutique, comprenant comme composant actif un composé tel que défini dans la revendication 1 ou 8, et un véhicule approprié pour celui-ci.
- 13. Utilisation d'un composé tel que défini dans la revendication 1, pour la fabrication d'un médicament ayant une activité d'antidepresseur et/ou soulageant un dysfonctionnement de la mémoire.
 - 14. Utilisation d'un composé tel que défini dans la revendication 8, pour la fabrication d'un médicament ayant une activité d'antidépresseur.
- 15. Procédé pour la préparation d'un composé tel que défini dans la revendication 1, comprenant
 - a) la réduction d'un composé de formule XVI

$$\begin{array}{c}
R_{12} \\
X
\end{array}$$

$$\begin{array}{c}
R_{3} \\
\end{array}$$

$$\begin{array}{c}
R_{3}
\end{array}$$

$$\begin{array}{c}
R_{3}
\end{array}$$

$$\begin{array}{c}
R_{3}
\end{array}$$

dans laquelle $\rm H_3$. X et n sont tels que définis dans la revendication 1, et $\rm R_{12}$ est un atome d'hydrogène ou le groupe méthoxy ou hydroxy, pour l'obtenition d'un composé de formule I dans lequel $\rm R_5$. X et n sont tels que définis, $\rm R_2$ est un atome d'hydrogène ou le groupe méthoxy ou hydroxy, et $\rm R_1$ et $\rm R_2$ représentent des atomes d'hydrogène ou

b) la mise en réaction d'un composé de formule XV

dans laquelle H_3 , X et n sont tels que définis dans la revendication 1, et H_{12} est un atome d'hydrogène ou le groupe méthoxy ou hydroxy, avec de l'isopropylate de titane et un composé de formule

dans laquelle le groupe

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$$-N$$
 NH $-N$ N-C₁-C₆-alkyle

$$-N$$
 N-phényle ou $-N$ $N-C_1\cdot C_5$ alkylphényle .

le fragment phényle pouvant être substitué comme indiqué dans la revendication 1.

suivie d'une réduction avec du borohydrure de sodium, pour la formation d'un composé de formule I dans lequel R_2 , X et n sont tels que définis, R_4 est tel que défini pour R_{10} ci-dessus, et le groupe

dans son ensemble a la signification donnée pour

ci-dessus, ou

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c) la réduction d'un composé de formule XVIII

$$\begin{array}{c} R_{11} \\ \\ X \\ \\ R_{1} \\ \\ \end{array}$$

dans laquelle R_3 . X et n sont tels que définis dans la revendication 1, R_{12} est un atome d'hydrogène ou le groupe méthoxy ou hydroxy, et R_1 est un groupe alkyle en C_1 - C_6 , alcényle en C_2 - C_6 , alcheryle en C_3 - C_6 , alcheryle en C_3 - C_6 , pricheryle-colakyle(C_3 - C_7) ou prényle, le fragment phényle pouvant être substitué comme indiqué dans la revendication 1, pour la formation d'un composé de formule I dans lequel R_3 . X et n sont tels que définis dans la revendication 1, R_{12} est tel que défini plus haut, R_2 est un atome d'hydrogène, et R_1 est tel que défini pour R_{14} ci-dessus,

d) éventuellement la réduction d'un composé de formule I dans lequel $R_{\rm S}$, $R_{\rm d}$, X et n sont tels que définis dans la revendication 1, et $R_{\rm l}$ et $P_{\rm S}$ représentent des atomes d'hydrogène, à l'aide de borane/feirahydrofuranne et d'acide trifluoroacétique, pour l'obtention d'un composé de formule la

dans laquelle R₃, R₄, X et n sont tels que définis,

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e) éventuellement la mise en réaction d'un composé de formule I dans lequel R_3 , R_4 , X et n sont tels que définis dans la revendication 1, et R_1 et R_2 sont des atomes d'hydrogène, avec un composé de formule Hal- R_{15} , dans laquelle R_{15} est un groupe alleyle en C_1 - C_6 , alcényle en C_2 - C_6 , alchynyle en C_3 - C_6 , cycloaikyl(C_7 - C_8), phényl-alkyle(C_1 - C_9), dans lequel le fragment phényle peut être substitué comme indiqué dans la revendication 1, ou un groupe de formule

Alk et Y étant tels que définis, pour l'obtention d'un composé de formule I dans lequel R_3 , R_4 , X et n sont tels que définis, R_1 a la signification de R_{15} telle que donnée plus haut, et R_2 est un atome d'hydrogène,

I) éventuellement la mise en réaction d'un composé de formule I dans lequel R_0 , R_0 , X at n sont tels que définis dans la revendication 1, R_1 , est un atome d'hydrogène ou un groupe allyle en C_2 - C_2 , cycloalityle en C_2 - C_2 , cycloalityle en C_2 - C_2 , prichapityle (C_1 - C_2) ou phényl-cycloalityle (C_2 - C_2), exprendent phényle pouvant être substitué comme indiqué dans la revendication 1, et R_1 , est un atome d'hydrogène, avec de l'acide formique, pour l'obtention d'un composé de formule I dans lequel R_3 , R_4 , X et n sont tels que définis dans la revendication 1, R_1 , est tel que défini plus haut et R_2 est le groupe formyle, g) eventuellement la mise en réaction d'un composé de formule I dans lequel R_3 , R_4 , X et n sont tels que défini dans la revendication 1, R_1 et un atome d'hydrogène ou un groupe altyle en C_2 - C_2 , elscriyle en C_2 - C_2 , cycloalityle en C_2 - C_3 , exprendent C_3 - C_3 , et C_3 - C_3 , exprendent C_3 - C_3 , exprendent C_3 - C_3

h) éventuellement la mise en réaction avec un chloroformiate de benzyle d'un composé de formule I dans lequel R_3 , R_4 , X et n sont tels que définis dans la revendication 1, étant entendu que R_4 n'est pas le groupe hydroxy, R_1 , est un atome d'hydrogène ou un groupe alkyle en $C_1 \cdot C_6$, alcényle en $C_2 \cdot C_6$, alcényle en $C_3 \cdot C_7$, phényl-alkyle($C_1 \cdot C_6$) ou phényl-cyclo- alkyle($C_3 \cdot C_7$), le fragment phényle pouvant être substitué comme indiqué dans la revendication 1, et R_2 est un atome d'hydrogène, pour l'obtention d'un composé de formule I dans lequel $R_3 \cdot R_4$, X et n sont tels que définis plus haut, R_1 est le groupe beavoyloxycarbonylox.

I) évantuellement la mise en réaction d'un composé de formule I dans la que IR $_0$, R $_4$, X et n sont tels que définis dans la revendication 1, étant entendu que R $_4$ n'est pas le groupe hydroxy, R $_1$ est un atome d'hydrogène ou un groupe alkyle en C $_1$ -C $_0$, alcényle en C $_2$ -C $_0$, alchyle en C $_3$ -C $_0$, cycloakyle en C $_3$ -C $_0$, cycloakyle en C $_3$ -C $_0$, prényl-alkyle (C $_3$ -C $_0$) ou phényl-eycloakyle (C $_3$ -C $_0$) et R $_0$ est un atome d'hydrogène, avec un isocyanate de formule R $_1$ -N-C $_0$ -C dans lequel R $_1$; est un groupe alkyle en C $_1$ -C $_0$, pényle ou phényl-alkyle (C $_1$ -C $_0$), le fragment phényle dans la définition de R $_1$ et R $_1$ 7 pouvant être substitué comme indiqué dans la revendication 1, pour l'obtention d'un composé de formule I dans lequel R $_3$, R $_4$, X et n sont tels que définis plus haut. R $_1$ est tel que défini plus haut, et R $_1$ est un groupe

dans lequel R₁₇ est tel que défini,

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j) éventuellement la mise en réaction d'un composé de formule I, dans lequel R_1 , R_2 , R_3 , X et n sont tels que définis dans la revendication 1, et R_1 est le groupe hydroxy, étant entendu que R_2 n'est pas un groupe alxyl $\{C_1,C_2\}$ amino-carbonyle, avec un chloroformiate de formule

dans laquelle R_7 est tel que défini dans la revendication 1, pour l'obtention d'un composé de formule I dans lequel R_1 , R_2 , R_3 , X et n sont tels que définis plus haut, et R_4 est un groupe

dans lequel R7 est tel que défini dans la revendication 1,

k) éventuellement la mise en réaction d'un composé de formule I dans lequel B_1 , B_3 , X et n sont tels que définis dans la revendication 1, B_2 est un atome d'hydrogène et B_4 est le groupe

dans lequel R₇ est le groupe benzyle, avec un isocyanate de formule R₁₇-N=C=O, pour l'obtention d'un composé de formule I dans lequel R₁, R₂, R₃, R₄, X et n sont tels que définis plus haut, et R₂ est un groupe

dans lequel R_{17} est un groupe alkyle en C_1 - C_6 , phényle ou phényl-alkyle (C_1 - C_6), le fragment phényle pouvant être substitué comme indiqué dans la revendication 1,

I) éventuellement la mise en réaction d'un composé de formule I dans lequel \mathbb{R}_1 , \mathbb{R}_2 , \mathbb{R}_3 \mathbb{X} et n sont tels que définis dans la revendication 1, et \mathbb{R}_4 est le groupe hydroxy, avec un composé de formule \mathbb{R}_1 - \mathbb{R}_2 - \mathbb{R}_2 - \mathbb{R}_3 - \mathbb{R}_4 -

dans lequel R₁₇ est tel que défini plus haut

16. Procédé pour la préparation d'un composé de formule III selon la revendication 8, comprenant

a) la mise en réaction d'un composé de formule XV

dans laquelle R_3 . X et n sont tels que définis dans la revendication 1 et R_{12} est un atome d'hydrogène ou le groupe hydroxy ou méthoxy, avec du chlorhydrate d'hydroxylamine, pour l'obtention d'un composé de formule III dans lequel R_3 , X et n sont tels que définis plus haut, R_4 a la signification de R_{12} donnée plus haut, et R_3 est le groupe hydroxy,

b) éventuellement la mise en réaction d'un composé de formule III dans lequel R_3 , X et n sont tels que définis dans la revendication 1, R_4 est un atome d'hydrogène ou le groupe hydroxy ou méthoxy, et R_6 est un atome d'hydrogène, avec un composé de formule $Br-R_13^*NH_2^1$, dans laquelle R_{13} est un groupe a lixylène en $C_1^*C_6$, pour l'obtention d'un composé de formule III dans lequel R_3 , R_{12} , X et n sont tels que définis et R_6 est un groupe aminoaloxy($C_1^*C_6$), $C_1^*C_6$.

c) la mise en réaction d'un composé de formule XV

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dans laquelle R_3 . X et n sont tels que définis dans la revendication 1 et R_{12} est un atome d'hydrogène ou le groupe hydroxy ou méthoxy, avec une amine de formule $Nt_2^4R_{14}$ dans laquelle R_{14} , et su n groupe alkyle en C_3 - C_3 , helfonyle en C_3 - $C_$

dans lequel R₁₇ est tel que défini,

 e) éventuellement la mise en réaction d'un composé de formule III dans lequel R₃. X et n sont tels que définis dans la revendication 1. R₄ est un atome d'hydrogène ou le groupe méthoxy ou hydroxy, et R₅ est le groupe hydroxy, avec un chlorure d'acyle de formule

ou un anhydride d'acide de formule $(R_{17}\cdot CO)_2O$ dans laquelle R_{17} est un groupe alkyle en $C_1\cdot C_8$ phényle ou phényl-alkyle $(C_1\cdot C_8)$, le fragment phényle pouvant être substitué comme indiqué dans la revendication 1, pour l'obtention d'un composé de formule III dans lequel R_9 , R_4 , X et n sont tels que définis plus haut et R_8 est le groupe

dans lequel R₁₇ est tel que défini plus haut.

17. Composé de formule II

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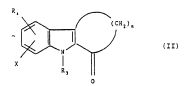
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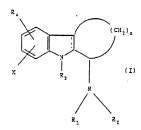
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dans laquelle R₃, R₄, X et n sont tels que définis dans la revendication 1.

35 Revendications pour les Etats contractants suivants : ES, GR

1. Procédé pour la préparation d'un composé de formule I



dans laquelle

n est 2, 3, 4 ou 5;

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- X est un atome d'hydrogène ou d'halogène ou un groupe alkyle en C₁-C₆, alcoxy en C₁-C₆, hydroxy, trifluorométhyle ou nitro;
- R₁ est un atome d'hydrogène ou un groupe allyte en C₁-C₆, alcényle en C₂-C₆, alcynyle en C₃-C₆, aminoalkyle en C₁-C₆, allyl(C₁-C₆), diallyl(C₁-C₆), cipclasityle en C₃-C₆, cycloalkyle en C₃-C₇, et cycloalkyle (C₂-C₇), le groupe phényle étant substitué par 0, 1 ou 2 substituants, représentant chacun indépendamment un atome d'halogène ou un groupe allèlyle en C₁-C₆, alcoys en C₁-C₆, lifturorméthyle, hydroxy ou nitro.

"alk" représentant un groupe alkylène divalent en C_1 - C_6 , et Y représentant un atome d'hydrogène ou un groupe alkylè en C_1 - C_6 , phényle ou phényl-alkyle(C_1 - C_6), le groupe phényle pouvant être substitué comme indiqué plus haut;

 R_2 est un atome d'hydrogène ou un groupe alkyle en C_1 - C_6 , formyle, alkyl(C_1 - C_6)-carbonyle, benzyloxycarbonyle ou alkyl(C_1 - C_6)-arbonyle, benzyloxycarbonyle, ou bien le groupe

dans son ensemble est

$$\frac{1}{2}$$
 $\frac{1}{2}$ $\frac{1}{2}$

$$-N \qquad NH \qquad -N \qquad N-C_1-C_6-alkyle$$

ou

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le radical phényle pouvant être substitué comme indiqué plus haut,

R₃ est un atome d'hydrogène ou un groupe alkyle en C₁-C₆, phényl-alkyle(C₁-C₆), le fragment phényle pouvant être substitué comme indiqué plus haut, alkyl(C₁-C₆)-carbonyle ou alcoxy(C₁-C₆)-carbonyle;

R₄ est un atome d'hydrogène, -OH, un groupe

ou

dans lequel

- R_{S} est un radical alkyle en $\mathsf{C}_1\mathsf{C}_{\mathsf{S}}$, alcónyle en $\mathsf{C}_2\mathsf{C}_{\mathsf{S}}$, alcynyle en $\mathsf{C}_3\mathsf{C}_{\mathsf{S}}$, cycloalkyle en $\mathsf{C}_3\mathsf{C}_{\mathsf{S}}$, cycloalkyle en $\mathsf{C}_3\mathsf{C}_{\mathsf{S}}$, cycloalkyle $(\mathsf{C}_2\mathsf{C}_{\mathsf{S}})$, alcónyle en $\mathsf{C}_3\mathsf{C}_{\mathsf{S}}$, cycloalkyle $(\mathsf{C}_3\mathsf{C}_{\mathsf{S}})$, le fragment phényle pouvant être substitué comme indiqué plus naut; et
- R₆ est un atome d'hydrogène ou un groupe alkyle en C₁-C₆, phényle ou phényl-alkyle(C₁-C₆), le fragment phényle pouvant être substitué comme indiqué plus haut; ou bien le groupe

est dans son ensemble

$$-N \qquad -N \qquad -N \qquad NH$$

$$-N \qquad N-C_1\cdot C_{\sigma} \text{-alkyle}$$

$$-N \qquad N-C_1\cdot C_{\sigma} \text{-alkylphānyle}.$$

le fragment phényle pouvant être substitué comme indiqué plus haut,

et

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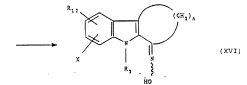
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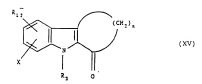
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- R₇ est un groupe alkyle en C₁-C₆, phényle ou phényl-alkyle(C₁-C₆), le fragment phényle pouvant être substitué comme indiqué plus haut;
- étant entendu que R₄ n'est pas un atome d'hydrogène ou le groupe hydroxy lorsque n est 4 ou 5; ou d'un sel d'addition avec un acide pharmaceutiquement acceptable de celui-ci, comprenant
 - a) la réduction d'un composé de formule XVI



dans laquelle R_3 , X et n sont tels que définis plus haut, et R_{12} est un atome d'hydrogène ou le groupe méthoxy ou hydroxy, pour l'obtention d'un composé de formule i dans lequel R_3 , X et n sont tels que définis, R_4 est un atome d'hydrogène ou le groupe méthoxy ou hydroxy, et R_1 et R_2 représentent des atomes d'hydrogène, ou b) la mise en réaction d'un composé de formule XV



dans laquelle H_3 , X et n sont tels que définis plus haut, et R_{12} est un atome d'hydrogène ou le groupe méthoxy ou hydroxy, avec de l'isopropylate de titane et un composé de formule

dans laquelle le groupe

est

$$-N$$
 NH $-N$ N-C₁-C₅-alkyle

50 le fragment phényle pouvant être substitué comme indiqué plus haut,

suivie d'une réduction avec du borohydrure de sodium, pour la formation d'un composé de formule I dans lequel R_a , X et n sont tels que définis, R_a est tel que défini pour R_{12} ci-dessus, et le groupe

dans son ensemble a la signification donnée pour



ci-dessus, ou

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c) la réduction d'un composé de formule XVIII

dans iaquelle R_3 . X et n sont tels que définis plus haut, R_{12} est un atorne d'hydrogène ou le groupe méthoxy ou hydroxy, et R_{14} est un groupe alkyle en C_3 - C_6 , alcényle en C_2 - C_6 , alcynyle en C_3 - C_7 , potényle, en C_3 - C_7 , potényle, le fragment phényle pouvant être substitué comme indiqué plus haut, pour la formation d'un composé de formule I dans lequel R_3 . X et n sont tels que définis plus haut, R_2 est tel que défini plus haut, R_2 est un atorne d'hydrogène, et R_1 est leque défini pour R_2 ci-dessus,

d) éventuellement la réduction d'un composé de formule I dans lequel H_3 , H_4 , X et n sont tels que définis plus haut, et H_1 et P_2 représentent des atomes d'hydrogène, à l'âide de borane/térahydrofuranne et d'acide tri-fluoroacétique, pour l'obtention d'un composé de formule la P_2

dans laquelle R₃, R₄, X et n sont tels que définis,

e) éventuellement la mise en réaction d'un composé de formule I dans lequel R₃, R₄, X et n sont tels que définisplus haut, et R₁ et R₂ sont des atomes d'hydrogène, avec un composé de formule Hal-R₁₅, dans laquelle R₁₅ est un groupe alkyle en C₁-C₆, alcényle en C₂-C₆, alcynyle en C₃-C₆, cycloalkyl(C₃-C₇), alcynyle en C₃-C₆, dans lequel le fragment phényle peut être substitué comme indiqué plus haut, ou un

groupe de formule

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Alk et Y étant tels que définis, pour l'obtention d'un composé de formule I dans lequel R_3 , R_4 , X et n sont tels que définis, R_1 a la signification de R_{15} telle que donnée plus haut, et R_2 est un atome d'hydrogène,

 $\hat{\bf 1}$) éventuellement la mise en réaction d'un composé de formule $\bf 1$ dans lequel $\bf R_3, R_4$. X et n sont tels que définis plus haut. $\bf R_1$ est un atome d'hydrogene ou un groupe alixyle en $\bf C_1$ - $\bf C_6$, alcényle en $\bf C_2$ - $\bf C_6$, alcynyle en $\bf C_3$ - $\bf C_7$, présyle en $\bf C_2$ - $\bf C_7$, présyle en $\bf C_3$ - $\bf C_7$, présyle pouvant être substitué comme indiqué plus haut, et $\bf R_2$ est un atome d'hydrogène, avec de l'acide formique, pour l'obtention d'un composé de formule I dans lequel $\bf R_3$, $\bf R_4$. X et n sont tels que définis plus haut et $\bf R_7$ est tiel que définis plus haut et $\bf R_7$ est tiel que définis plus haut et $\bf R_7$ est tiel que définis plus haut et $\bf R_7$ est tiel que définis plus haut et $\bf R_7$ est le que définis plus haut et $\bf R_7$ est le que que définis plus haut et $\bf R_7$ est le que définis plus haut et $\bf R_7$ est le que définis plus haut et $\bf R_7$ est le que définis plus haut et $\bf R_7$ est le que définis plus haut et $\bf R_7$ est le que définis plus haut et $\bf R_7$ est le que définis plus haut et $\bf R_7$ est et que définis plus haut et $\bf R_7$ est et que définis plus haut et $\bf R_7$ est et que définis plus haut et $\bf R_7$ est et que définis plus haut et $\bf R_7$ est et que définis plus haut et $\bf R_7$ est et que définis plus haut et $\bf R_7$ est et que définis plus haut et $\bf R_7$ est et que de l'aux et $\bf R_7$ est et que de l'aux et $\bf R_7$ est et $\bf R_7$ e

g) éventuellement la mise en réaction d'un composé de formule I dans lequel R_3 , R_4 . X et n sont tels que définis plus haut, R_1 et un atome d'hydrogène ou un groupe alkyle en C_1 - C_6 , alcènyle en C_2 - C_6 , alcènyle en C_3 , alcènyle

h) èventuellement la mise en réaction avec un chloroformiste de benzyle d'un composé de formule I dans lequel Π_{S_1} , Π_{A_1} X et n sont tels que définis, étant entant que Π_{A_1} et pas le groupe $hypricxy_1$, Π_{A_1} est un atomytrogène ou un groupe altèlyle en $G_1 G_2$, alcényle en $G_2 G_3$, alcynyle en $G_2 G_3$, cycloalistyle en $G_2 G_3$, cycloalistyle en $G_2 G_3$, phényl-alikyle $(G_1 G_2)$ ou phényl-cycloalisyle $(G_2 G_3)$, le fragment phényle pouvant être substituté comme indiqué plus haut, et Π_{S_2} et un atome d'hydrogène, pour l'obtention d'un composé de formule I dans lequel Π_{S_2} Π_{S_3} X et n sont tels que définis. Π_{S_3} est tel que défini plus haut, et Π_{S_2} est le groupe benzy-loxycarbonyle.

i) éventuellement la mise en réaction d'un composé de formule I dans lequel R_b, R_b, X ant sont tels que définis plus haut, étant entendu que R_b n'est pas le groupe hydroxy, R_1 est un atome d'hydrogène ou un groupe alkyle en $C_3 - C_6$, coloakényle en $C_3 - C_6$, alcynyle en $C_3 - C_6$, alcynyle en $C_3 - C_6$, coloakényle en $C_3 - C_6$, prohayl-alkyle $(C_1 - C_6)$ ou phényl-cyclealkyle $(C_3 - C_7)$. le fragment phényle pouvant être substitué comme indiqué plus haut, et R_3 est un entom d'hydrogène, avec un isocyanate de formule $R_1 - N - C_6$, chényle ou phényl-alkyle $(C_1 - C_6)$, le fragment phényle pouvant être substitué comme indiqué plus haut, pour l'obtention d'un composé de formule I dans lequel R_3 , R_4 , X et n sont tels que définis, R_1 est le que défini plus haut,

et R2 est un groupe

dans lequel R₁₇ est tel que défini,

 j) éventuellement la mise en réaction d'un composé de formule I, dans lequel R₁, R₂, R₃, X et n sont tels que définis, et R₄ est le groupe hydroxy, étant entendu que R₂ n'est pas un groupe alkyl(C₁-C₆)amino-carbonyle, avec un chloroformiate de formule

dans laquelle R_7 est tel que défini, pour l'obtention d'un composé de formule I dans lequel R_1 , R_2 , R_3 , X et n sont tels que définis, et R_4 est un groupe

dans lequel R7 est tel que défini,

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k) éventuellement la mise en réaction d'un composé de formule I dans lequel R_1 , R_3 . X et n sont tels que définis, R_2 est un atome d'hydrogène et R_4 est le groupe

dans lequel R₇ est le groupe benzvle.

avec un isocyanate de formule R_{17} -N=C=O, pour l'obtention d'un composé de formule I dans lequel R_1 , R_2 , R_3 , R_4 , X et n sont tels que définis, et R_5 est un groupe

dans lequel R_{17} est un groupe alkyle en C_1 - C_6 , phényle ou phényl-alkyle(C_1 - C_6), le fragment phényle pouvant être substitué comme indiqué plus haut,

I) éventuellement la mise en réaction d'un composé de formule I dans lequel R_1 , R_2 , R_3 , X et n sont tels que définis, et R_4 est le groupe hydroxy, avec un composé de formule R_{17} -NCCO dans lequel R_{17} est un groupe alkyle en C_1 - C_6 , phényle ou phényl-alkyle $(C_1$ - C_6), le fragment phényle pouvant être substitué comme indiqué plus haut, pour l'obtention d'un composé de formule I dans lequel R_1 , R_2 , R_3 X et n sont tels que définis plus haut, et R_3 est un groupe

dans lequel R₁₇ est tel que défini plus haut.

- 2. Procédé selon la revendication 1, dans lequel n est égal à 3.
- 3. Procédé selon la revendication 2, dans lequel
 - X est un atome d'hydrogène ou le groupe hydroxy,
 - R₁ est un atome d'hydrogène ou un groupe alcynyle en C₃-C₆, cycloalkyle en C₃-C₇, phényle ou phényl-alkyle
 - R₂ est un atome d'hydrogène ou un groupe formyle, benzyloxycarbonyle ou alkyl(C₁-C₆)amino-carbonyle,
 - R3 est un atome d'hydrogène ou un groupe alkyle en C1-C6
 - R4 est un atome d'hydrogène ou un groupe de formule

formules dans lesquelles R_g est un groupe alkyle en C_1 – C_g ou phényl-alkyle(C_1 – C_g) et R_g est un atome d'hydrogène, et R_p est un groupe phényl-alkyle(C_1 – C_g), chaque groupe phényle dans les définitions de R_1 , R_g et R_g pouvant être substitué comme indiqué dans la revendication 1.

- 4. Procédé selon la revendication 3. dans lequel
 - X est un atome d'hydrogène.

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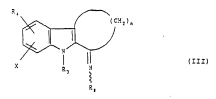
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- R₁ est un groupe cycloaikyle en C₃-C₇, alcynyle en C₃-C₆, phényl-cycloaikyle(C₃-C₇) ou phényl-alkyle(C₁-C₆), le fragment phényle pouvant être substitué comme indiqué dans la revendication 1,
- R₂ est un atome d'hydrogène,
- R4 est un atome d'hydrogène ou un groupe de formule

dans laquelle R₅ est un groupe alkyle en C₁-C₆ et R₆ est un atome d'hydrogène.

- Procédé selon la revendication 1, dans lequel on prépare le méthylcarbonate de 3-cyclopropylamino-4-méthyl-1,2,3,4-tétrahydrocyclopent[b]indole-7-yle.
- Procédé selon la revendication 1, dans lequel on prépare le méthylcarbonate de 4-méthyl-3-phénylméthylamino-1,2,3,4-tétrahydrocyclopent[b]indole-7-yle.
 - Procédé selon la revendication 1, dans lequel on prépare la 1,2,3,4-tétrahydrocyclopent[b]indole-3-(2-propynyl)
 amine.
 - 8. Procédé pour la préparation d'un composé de formule III



dans laquelle R_3 , R_4 . X et n sont tels que définis dans la revendication 1, et R_6 est un groupe hydroxy, alcoxy en $C_1 \cdot C_6$, aiminoalcoxy($C_1 \cdot C_6$), alityle en $C_1 \cdot C_6$, alcyryle en $C_3 \cdot C_6$, cycloalkyle en $C_3 \cdot C_6$, cycloalkyle en $C_3 \cdot C_6$, cycloalkyle en $C_3 \cdot C_6$, phényladyleighe($C_1 \cdot C_6$) ou phényl-cycloalkyle($C_3 \cdot C_6$), le fragment phényle pouvant être substituté comme indiqué dans la revendication 1, alkyle($C_1 \cdot C_6$), acrahomyloxy ou alkyl($C_1 \cdot C_6$) aminocarbonyloxy, ou d'un sel d'addition pharmaceutiquement acceptable de celui-ci, comprenant

a) la mise en réaction d'un composé de formule XV

dans laquelle R_3 . X et n sont tels que définis dans la revendication 1 et R12 est un atome d'hydrogène ou le groupe hydroxy ou méthoxy, avec du chlorhydrate d'hydroxylamine, pour l'obtention d'un composé de formule III dans lequel R_3 . X et n sont tels que définis plus haut, R_4 a la signification de R_{12} donnée plus haut, et R_3 est le groupe hydroxy.

b) eventuellement la mise en réaction d'un composé de formule III dans lequel R_b . X et n sont tels que définis dans la revendication 1, R_b est un atome d'hydrogène ou le groupe hydroxy ou méthoxy, et R_b est un atome d'hydrogène, avec un composé de formule R_b , R_b - $R_$

c) la mise en réaction d'un composé de formule XV

dans laquelle P_0 , X et n sont tels que définis dans la revendication 1 et P_0 , est un atome d'hydrogène ou le groupe hydroy ou méthoxy, avec une amine de formule NH₂P₁, dans laquelle P_0 , est un groupe alkyle en $C_3 \cdot C_6$, alcényle en $C_2 \cdot C_6$, alcényle en $C_3 \cdot C_6$, opcloalkyle en $C_3 \cdot C_7$, cycloalcényle en $C_3 \cdot C_7$, phényl-alkyle($C_1 \cdot C_6$), phényl-cycloalkyle $(C_3 \cdot C_6)$, op hényl-alkyle($C_3 \cdot C_6$), operation of the compose de formule III dans lequel P_0 , X et n sont lest que définis plus haut. P_0 a la signification de P_1 , donnée plus haut, et P_0 a la signification de P_1 , donnée plus haut, et P_0 a la signification de P_1 , donnée plus haut, et P_0 et entre les que définis dans la revendication P_0 exception de P_1 , donnée plus haut, et P_0 exception P_0 exception

dans lequel R₁₇ est tel que défini,

e) éventuellement la mise en réaction d'un composé de formule III dans lequel R_3 . X et n sont tels que définis dans la revendication 1. R_4 est un atome d'hydrogène ou le groupe méthoxy ou hydroxy, et R_3 est le groupe hydroxy, avec un chlorure d'acyle de formule

ou un anhydride d'acide de formule $(R_{17}\cdot CO)_2O$ dans laquelle R_{17} est un groupe alkyle en $C_1\cdot C_6$ phényle ou phényl-alkyle $(C_1\cdot C_6)$, le fragment phényle pouvant être substitué comme indiqué dans la revendication 1, pour l'obtention d'un composé de formule III dans lequel R_9 . R_4 . X et n sont tels que définis plus haut et R_6 est le groupe

dans lequel R₁₇ est tel que défini plus haut.

- 9. Procédé selon la revendication 8, dans lequel n est égal à 3.
- 20 10. Procédé selon la revendication 9, dans lequel

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- X est un atome d'hydrogène ou un groupe hydroxy ou alcoxy en C₁-C₂.
- R₃ est un atome d'hydrogène ou un groupe alkyle en C₁-C₆,
- R4 est un atome d'hydrogène ou un groupe de formule

formules dans lesquelles $\rm H_5$ est un groupe alkyle en $\rm C_1$ - $\rm C_6$ ou phényl-alkyle($\rm C_1$ - $\rm C_6$), $\rm H_6$ est un atome d'hydrogène. et $\rm H_7$ est un groupe alkyle en $\rm C_1$ - $\rm C_6$ ou phényl-alkyle($\rm C_1$ - $\rm C_6$),

- R₈ est un groupe hydroxy, akcynyle en C₃-C₆, amino-alcoxy(C₁-C₆), alklyl(C₁-C₆)-carbonyloxy, alklyl(C₁-C₆)-carbonyloxy, alklyl(C₁-C₆)-carbonyloxy, alklyl(C₁-C₆), chaque fragment phényje dans les définitions de R₅. R₅ et R₅ pouvant être substitué comme indiqué dans la revendication 1.
- Procédé selon la revendication 8, dans lequel on prépare le 4-méthyl-3-phénylméthylimino-1,2,3,4-tétrahydrocyclopent[b]indole-7-ol.
- 12. Utilisation d'un composé tel que défini dans la revendication 1, pour la fabrication d'un médicament ayant une activité d'antidépresseur et/ou soulageant un dysfonctionnement de la mémoire.
- Utilisation d'un composé tel que défini dans la revendication 8, pour la fabrication d'un médicament ayant une activité d'antidépresseur.
- 14. Composé de formule II

dans laquelle H_3 , H_4 , X et n sont tels que définis dans la revendication 1.